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(54) Title: METHOD AND NUCLEIC ACIDS FOR THE ANALYSIS OF COLON CANCER

(57) Abstract: The present invention relates to chemically modified genomic sequences, oligonucleotides and/or PNA-oligomers for detecting the cytosine methylation state of genomic DNA, as well as to methods for ascertaining genetic and/or epigenetic parameters of genes for use in the characterisation, grading, staging, and/or diagnosis of colon cancer, or the predisposition to colon cancer.

## Method and nucleic acids for the analysis of colon cancer

### Field of the Invention

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The levels of observation that have been studied by the methodological developments of recent years in molecular biology, are the genes themselves, the translation of these genes into RNA, and the resulting proteins. The question of which gene is switched on at which point in the course of the development of an individual, and how the activation and inhibition of specific genes in specific cells and tissues are controlled is correlatable to the degree and character of the methylation of the genes or of the genome. In this respect, pathogenic conditions may manifest themselves in a changed methylation pattern of individual genes or of the genome.

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The present invention relates to nucleic acids, oligonucleotides, PNA-oligomers, and to a method for the characterisation, grading, staging, treatment and/or diagnosis of colon cancer, or the predisposition to colon cancer, by analysis of the genetic and/or epigenetic parameters of genomic DNA and, in particular, with the cytosine methylation status thereof.

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### Prior Art

Colon cancer is the second most common cause of cancer death in the United States. It describes any cancer in the colon (large intestine), from the beginning of the colon (cecum) to the end of the colon (rectum). Colon cancer is a malignant tumor in the lining of the large intestine. It starts with a single cell that mutates and grows into a visible polyp. If a polyp is allowed to remain in the colon it can grow into a cancerous tumor that

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can invade other organs. The mechanism behind the progression to malignancy are not completely understood, however most polyps take 3-7 years to become cancerous. Prevention of colon cancer means stopping this process by removing the polyp before it becomes cancerous. Colon cancer represents an interaction between the genome of the colorectal epithelial cell and the host environment. Both factors are essential for the development of tumors. Colon cancers can be differentiated into nonhereditary types, which rarely occur before age 40 and hereditary colon cancers which often occur in younger people.

Human colon cancers undergo a multistage carcinogenesis pathway from adenomatous polyps to carcinoma. A number of genetic events have been characterized and include alterations in "tumor suppressor" and susceptibility genes that normally encode for proteins regulating cell cycle progression and programmed cell death (Kinzler KW, Vogelstein B. Landscaping the cancer terrain. Science. 1998 May 15;280(5366):1036-7). Given the high incidence of colon cancer in the aging population and high mortality rates for advanced disease, new prevention strategies are needed. After the diagnosis of cancer has been made it is important to determine the extent or 'stage' of the cancer before deciding on the treatment plan. Staging is a method of evaluating the progress of the cancer in a patient and defines the extent to which the cancer has spread to other parts of the body. There are several systems for classifying the extent or stage of cancer. One of the the two most common systems is the Stage 'I, II, III, IV' system, which defines four stages of cancer. Stage I represents early cancer, with a small tumor and no spread to the lymph nodes. In stages II and III, the tumor is progressively more advanced, while stage IV refers to metastatic disease that has spread to other areas of the body. One very important point to realize about

these staging systems is that they only provide rough estimates of the stage of disease and chances of survival. The numbers are just averages. They do not say anything about the outcome or prognosis of any one particular patient.

Genes which are associated with colon cancer include the following.

p16 (Dai CY, Furth EE, Mick R, Koh J, Takayama T, Niitsu Y, Enders GH. p16(INK4a) expression begins early in human colon neoplasia and correlates inversely with markers of cell proliferation. Gastroenterology. 2000 Oct; 119(4):929-42).

p27 (Liu DF, Ferguson K, Cooper GS, Grady WM, Willis J. p27 cell-cycle inhibitor is inversely correlated with lymph node metastases in right-sided colon cancer. J Clin Lab Anal. 1999;13(6):291-5).

p53 (Arango D, Corner GA, Wadler S, Catalano PJ, Augenlicht LH. c-myc/p53 interaction determines sensitivity of human colon carcinoma cells to 5-fluorouracil in vitro and in vivo. Cancer Res. 2001 Jun 15;61(12):4910-5).  
cdc2 (Moragoda L, Jaszewski R, Majumdar AP. Curcumin induced modulation of cell cycle and apoptosis in gastric and colon cancer cells. Anticancer Res. 2001 Mar-Apr; 21(2A):873-8).

PCNA (Zhang Y, Iwama T, Sugihara K. Histochemical study of apoptosis and cell proliferation in hereditary intestinal diseases. J Med Dent Sci. 1998 Jun;45(2):77-84).  
CEA (Vogel I, Francksen H, Soeth E, Henne-Bruns D, Kremer B, Juhl H. The carcinoembryonic antigen and its prognostic impact on immunocytologically detected intraperitoneal colorectal cancer cells. Am J Surg. 2001 Feb;181(2):188-93).

c-erbB2 (Fric P, Sovova V, Sloncova E, Lojda Z, Jirasek A, Cermak J. Different expression of some molecular mark-



ers in sporadic cancer of the left and right colon. *Eur J Cancer Prev.* 2000 Aug; 9(4):265-8).

Estrogen receptor (Campbell-Thompson M, Lynch IJ, Bhardwaj B. Expression of estrogen receptor (ER) subtypes and ERbeta isoforms in colon cancer. *Cancer Res.* 2001 Jan 15;61(2):632-40).

Progesterone receptor (Reich O, Regauer S, Urdl W, Lahousen M, Winter R. Expression of oestrogen and progesterone receptors in low-grade endometrial stromal sarcomas. *Br J Cancer.* 2000 Mar;82(5):1030-4) and myoglobin (Nakao A, Sakagami K, Uda M, Mitsuoka S, Ito H. Carcinosarcoma of the colon: report of a case and review of the literature. *J Gastroenterol.* 1998 Apr;33(2):276-9).

5-methylcytosine is the most frequent covalent base modification in the DNA of eukaryotic cells. It plays a role, for example, in the regulation of the transcription, in genetic imprinting, and in tumorigenesis. Therefore, the identification of 5-methylcytosine as a component of genetic information is of considerable interest. However, 5-methylcytosine positions cannot be identified by sequencing since 5-methylcytosine has the same base pairing behavior as cytosine. Moreover, the epigenetic information carried by 5-methylcytosine is completely lost during PCR amplification.

A relatively new and currently the most frequently used method for analyzing DNA for 5-methylcytosine is based upon the specific reaction of bisulfite with cytosine which, upon subsequent alkaline hydrolysis, is converted to uracil which corresponds to thymidine in its base pairing behavior. However, 5-methylcytosine remains unmodified under these conditions. Consequently, the original DNA is converted in such a manner that methylcytosine, which originally could not be distinguished from cytosine by its hybridization behavior, can now be de-

5        tected as the only remaining cytosine using "normal" molecular biological techniques, for example, by amplification and hybridization or sequencing. All of these techniques are based on base pairing which can now be fully  
10        exploited. In terms of sensitivity, the prior art is defined by a method which encloses the DNA to be analyzed in an agarose matrix, thus preventing the diffusion and renaturation of the DNA (bisulfite only reacts with single-stranded DNA), and which replaces all precipitation  
15        and purification steps with fast dialysis (Olek A, Oswald J, Walter J. A modified and improved method for bisulfite based cytosine methylation analysis. Nucleic Acids Res. 1996 Dec 15;24(24):5064-6). Using this method, it is possible to analyze individual cells, which illustrates  
20        the potential of the method. However, currently only individual regions of a length of up to approximately 3000 base pairs are analyzed, a global analysis of cells for thousands of possible methylation events is not possible. However, this method cannot reliably analyze very small  
25        fragments from small sample quantities either. These are lost through the matrix in spite of the diffusion protection.

25        An overview of the further known methods of detecting 5-methylcytosine may be gathered from the following review article: Rein, T., DePamphilis, M. L., Zorbas, H., Nucleic Acids Res. 1998, 26, 2255.

30        To date, barring few exceptions (e.g., Zeschnigk M, Lich C, Buiting K, Doerfler W, Horsthemke B. A single-tube PCR test for the diagnosis of Angelman and Prader-Willi syndrome based on allelic methylation differences at the SNRPN locus. Eur J Hum Genet. 1997 Mar-Apr;5(2):94-8) the bisulfite technique is only used in research. Always,  
35        however, short, specific fragments of a known gene are amplified subsequent to a bisulfite treatment and either

- completely sequenced (Olek A, Walter J. The pre-implantation ontogeny of the H19 methylation imprint. *Nat Genet.* 1997 Nov;17(3):275-6) or individual cytosine positions are detected by a primer extension reaction (Gonzalzo ML, Jones PA. Rapid quantitation of methylation differences at specific sites using methylation-sensitive single nucleotide primer extension (Ms-SNuPE). *Nucleic Acids Res.* 1997 Jun 15;25(12):2529-31, WO Patent 9500669) or by enzymatic digestion (Xiong Z, Laird PW. COBRA: a sensitive and quantitative DNA methylation assay. *Nucleic Acids Res.* 1997 Jun 15;25(12):2532-4). In addition, detection by hybridization has also been described (Olek et al., WO 99 28498).
- Further publications dealing with the use of the bisulfite technique for methylation detection in individual genes are: Grigg G, Clark S. Sequencing 5-methylcytosine residues in genomic DNA. *Bioessays.* 1994 Jun;16(6):431-6, 431; Zeschmick M, Schmitz B, Dittrich B, Buiting K, Horsthemke B, Doerfler W. Imprinted segments in the human genome: different DNA methylation patterns in the Prader-Willi/Angelman syndrome region as determined by the genomic sequencing method. *Hum Mol Genet.* 1997 Mar;6(3):387-95; Feil R, Charlton J, Bird AP, Walter J, Reik W. Methylation analysis on individual chromosomes: improved protocol for bisulphite genomic sequencing. *Nucleic Acids Res.* 1994 Feb 25;22(4):695-6; Martin V, Ribieras S, Song-Wang X, Rio MC, Dante R. Genomic sequencing indicates a correlation between DNA hypomethylation in the 5' region of the pS2 gene and its expression in human breast cancer cell lines. *Gene.* 1995 May 19;157(1-2):261-4; WO 97/46705, WO 95/15373 and WO 95/45560.
- An overview of the Prior Art in oligomer array manufacturing can be gathered from a special edition of *Nature*

Genetics (Nature Genetics Supplement, Volume 21, January 1999), published in January 1999, and from the literature cited therein.

5 Fluorescently labeled probes are often used for the scanning of immobilized DNA arrays. The simple attachment of Cy3 and Cy5 dyes to the 5'-OH of the specific probe are particularly suitable for fluorescence labels. The detection of the fluorescence of the hybridized probes may be  
10 carried out, for example via a confocal microscope. Cy3 and Cy5 dyes, besides many others, are commercially available.

Matrix Assisted Laser Desorption Ionization Mass Spectrometry (MALDI-TOF) is a very efficient development for  
15 the analysis of biomolecules (Karas M, Hillenkamp F. Laser desorption ionization of proteins with molecular masses exceeding 10,000 daltons. Anal Chem. 1988 Oct 15;60(20):2299-301). An analyte is embedded in a light-absorbing matrix. The matrix is evaporated by a short laser pulse thus transporting the analyte molecule into the  
20 vapor phase in an unfragmented manner. The analyte is ionized by collisions with matrix molecules. An applied voltage accelerates the ions into a field-free flight tube. Due to their different masses, the ions are accelerated at different rates. Smaller ions reach the detector  
25 sooner than bigger ones.

MALDI-TOF spectrometry is excellently suited to the  
30 analysis of peptides and proteins. The analysis of nucleic acids is somewhat more difficult (Gut I G, Beck S. DNA and Matrix Assisted Laser Desorption Ionization Mass Spectrometry. Current Innovations and Future Trends. 1995, 1; 147-57). The sensitivity to nucleic acids is approximately 100 times worse than to peptides and  
35 decreases disproportionally with increasing fragment size.

For nucleic acids having a multiply negatively charged backbone, the ionization process via the matrix is considerably less efficient. In MALDI-TOF spectrometry, the selection of the matrix plays an eminently important  
5 role. For the desorption of peptides, several very efficient matrixes have been found which produce a very fine crystallization. There are now several responsive matrixes for DNA, however, the difference in sensitivity has not been reduced. The difference in sensitivity can  
10 be reduced by chemically modifying the DNA in such a manner that it becomes more similar to a peptide. Phosphorothioate nucleic acids in which the usual phosphates of the backbone are substituted with thiophosphates can be converted into a charge-neutral DNA using simple alkylation chemistry (Gut IG, Beck S. A procedure for selective DNA alkylation and detection by mass spectrometry. Nucleic Acids Res. 1995 Apr 25;23(8):1367-73). The coupling of a charge tag to this modified DNA results in an increase in sensitivity to the same level as that found  
15 for peptides. A further advantage of charge tagging is the increased stability of the analysis against impurities which make the detection of unmodified substrates considerably more difficult.

25 Genomic DNA is obtained from DNA of cell, tissue or other test samples using standard methods. This standard methodology is found in references such as Fritsch and Maniatis eds., Molecular Cloning: A Laboratory Manual, 1989.

### 30 Description of the invention

The present invention discloses that atypical methylation in the genes estrogen receptor, p21, p27, p16, progesteron receptor, myoglobin, pcna, cdc2, c-erbB2, p53  
35 and CEA, can be positively correlated with colon carcinogenesis. This allows the detection of colon carcinoma, or

the predisposition to colon cancer by an assay that detects methylation in the genes by restriction enzyme analysis, or using a nucleic acid based method.

5 The disclosed invention provides a method and nucleic acids for the analysis of colon carcinomas. It discloses a means of distinguishing between healthy and cancerous colon tissue. This provides a means for the improved diagnosis, prognosis, staging and grading of colon cancer, at  
10 a molecular level, as opposed to currently used methods of a relatively subjective nature such as histological analysis. Furthermore, the disclosed invention presents improvements over the state of the art in that current methods of histological and cytological analysis require  
15 that the biopsy contain a sufficient amount of tissue. The method according to the present invention can be used for classification of minute samples.

The invention provides a method for detecting a colon  
20 cell proliferative disorder characterised in that the target nucleic acid of one or more genes taken from the group comprising estrogen receptor, p21, p27, p16, progesteron receptor, myoglobin, pcna, cdc2, c-erbB2, p53 and CEA are contacted with a reagent or series of re-  
25 agents capable of distinguishing between methylated and non methylated CpG dinucleotides within the target sequence.

The present invention makes available a method for ascer-  
30 taining genetic and/or epigenetic parameters of genomic DNA. The method is for use in the grading, staging, treatment and/or diagnosis of colon cancer. The method enables the analysis of cytosine methylations and single nucleotide polymorphisms.

In one embodiment of the method the genomic DNA sample is first isolated from tissue or cellular sources. Such sources may include cell lines, histological slides, body fluids, or tissue embedded in paraffin. Extraction may be by means that are standard to one skilled in the art, these include the use of detergent lysates, sonification and vortexing with glass beads. Once the nucleic acids have been extracted the genomic double stranded DNA is used in the analysis.

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In a preferred embodiment the DNA may be cleaved prior to the chemical treatment, this may be any means standard in the state of the art, in particular with restriction endonucleases.

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In the third step of the method, the genomic DNA sample is treated in such a manner that cytosine bases which are unmethylated at the 5'-position are converted to uracil, thymine, or another base which is dissimilar to cytosine in terms of hybridization behavior. This will be understood as 'pretreatment' hereinafter.

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The above described treatment of genomic DNA is preferably carried out with bisulfite (sulfite, disulfite) and subsequent alkaline hydrolysis which results in the conversion of non-methylated cytosine nucleobases to uracil or to another base which is dissimilar to cytosine in terms of base pairing behavior.

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In the fourth step of the method the bisulfite treated DNA is analysed using one or a combination of several methods which are known in the art namely real time PCR (Methyl Light assay), blocking oligonucleotides, methylation specific single nucleotide polymorphism extension (hereinafter referred to as MsSNuPE), methylation spe-

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cific PCR (hereinafter referred to as MSP), and nucleic acid sequencing.

Fluorescence-based Real Time Quantitative PCR (Heid et al., Genome Res. 6:986-994, 1996) employs a dual-labeled fluorescent oligonucleotide probe (e.g. TaqMan™ PCR, using an ABI Prism 7700 Sequence Detection System, Perkin Elmer Applied Biosystems, Foster City, California) that is hybridized concurrently with oligonucleotide primers during a continuously monitored polymerase chain reaction.. The TaqMan™ PCR reaction employs the use of a nonextendible interrogating oligonucleotide, called a TaqMan™ probe, which is designed to hybridize to a GpC-rich sequence located between the forward and reverse amplification primers. The TaqMan™ probe further comprises a fluorescent "reporter moiety" and a "quencher moiety" covalently bound to linker moieties (e.g., phosphoramidites) attached to the nucleotides of the TaqMan™ oligonucleotide. For analysis of methylation within nucleic acids subsequent to bisulphite treatment it is required that the probe be methylation specific, as described in U.S. 6,331,393, also known as the Methyl Light assay. Variations on the TaqMan™ detection methodology that are also suitable for use with the described invention include the use of dual probe technology (Lightcycler™) or fluorescent amplification primers (Sunrise™ technology). Both these techniques may be adapted in a manner suitable for use with bisulphite treated DNA, and moreover for methylation analysis within CpG dinucleotides.

A further suitable method for the for the assessment of methylation by analysis of bisulphite treated nucleic acids is the use of blocker oligonucleotides. The use of such oligonucleotides has been described in BioTechniques 23(4), 1997, 714-720 D. Yu, M.Mukai, Q. Liu, C. Steinman. Blocking probe oligonucleotides are hybridised to the bi-



5 sulphite treated nucleic acid concurrently with the PCR primers. PCR amplification of the nucleic acid is terminated at the 5' position of the blocking probe, thereby amplification of a nucleic acid is suppressed wherein the complementary sequence to the blocking probe is present. The probes may be designed to hybridise to the bisulphite treated nucleic acid in a methylation status specific manner. For example, for detection of methylated nucleic acids within a population of unmethylated nucleic acids suppression of the amplification of nucleic acids which are unmethylated at the position in question would be carried out by the use of blocking probes comprising a 'CG' at the position in question, as opposed to a 'CA'.

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15 In a further preferred embodiment of the method the analysis is carried out by the use of template directed oligonucleotide extension, such as MS SNUPE as described by Gonzalzo and Jones (Nucleic Acids Res. 25:2529-2531).

20 In an alternative embodiment of the method the assessment of the methylation state for the CpG dinucleotides may be carried out by PCR analysis of the treated nucleic acid(s) using methylation specific PCR. Methylation specific primers (MSP) have been described, for example in

25 U.S. Patent 6,265,171 to Herman et al. MSP primers consist of an oligonucleotide specific for annealing to a nucleotide sequence containing at least one bisulphite treated CpG dinucleotide. Therefore the sequence of said primers includes at least one CG, TG or CA dinucleotide.

30 MSP primers specific for non methylated DNA contain a 'T' at the 3' position of the C position in the CpG. MSP primers generally contain relatively few cytosines as these are converted by the bisulphite reaction. However when the primers are specific for methylated cytosine dinucleotides said cytosine positions are conserved within

35 the primer oligonucleotides.

The primers are extended by means of a polymerase and the resultant double stranded nucleic is denatured, preferably by means of heat treatment. Successive cycles of primer annealing, extension and denaturation are carried out according to the polymerase chain reaction as described in U.S. Pat. No. 4,582,788 to Mullis.

In a further embodiment of the method the analysis is enabled by sequencing and subsequent sequence analysis of the amplificate generated in the third step of the method (Sanger F., et al., 1977 PNAS USA 74: 5463-5467).

In a particularly preferred embodiment, the method comprises the following steps:

In the first step of the method the genomic DNA sample must be isolated from tissue or cellular sources. Such sources may include cell lines, histological slides, body fluids, or tissue embedded in paraffin. Extraction may be by means that are standard to one skilled in the art, these include the use of detergent lysates, sonification and vortexing with glass beads. Once the nucleic acids have been extracted the genomic double stranded DNA is used in the analysis.

In a preferred embodiment the DNA may be cleaved prior to the chemical treatment, this may be any means standard in the state of the art, in particular with restriction endonucleases.

In the second step of the method, the genomic DNA sample is treated in such a manner that cytosine bases which are unmethylated at the 5'-position are converted to uracil, thymine, or another base which is dissimilar to cytosine

in terms of hybridization behavior. This will be understood as ' pretreatment' hereinafter.

5 The above described treatment of genomic DNA is preferably carried out with bisulfite (sulfite, disulfite) and subsequent alkaline hydrolysis which results in the conversion of non-methylated cytosine nucleobases to uracil or to another base which is dissimilar to cytosine in terms of base pairing behavior.

10 In the third step fragments of the pretreated DNA are amplified, using sets of primer oligonucleotides according to Seq ID 76 to 97, and a, preferably heat-stable polymerase. Because of statistical and practical considerations, preferably more than ten different fragments having a length of 100 - 2000 base pairs are amplified. The amplification of several DNA segments can be carried out simultaneously in one and the same reaction vessel. Usually, the amplification is carried out by means of a polymerase chain reaction (PCR).

20 The method may also be enabled by the use of alternative primers, the design of such primers is obvious to one skilled in the art. These should include at least two oligonucleotides whose sequences are each reverse complementary or identical to an at least 18 base-pair long segment of the base sequences specified in the appendix (Seq. ID No.32 through Seq. ID No.75). Said primer oligonucleotides are preferably characterized in that they do not contain any CpG dinucleotides. In a particularly preferred embodiment of the method, the sequence of said primer oligonucleotides are designed so as to selectively anneal to and amplify, only the colon tissue specific DNA of interest, thereby minimizing the amplification of background or non relevant DNA. In the context of the present invention, background DNA is taken to mean ge-

nomie DNA which does not have a relevant tissue specific methylation pattern, in this case, the relevant tissue being colon tissue, both healthy and diseased.

5 According to the present invention, it is preferred that at least one primer oligonucleotide is bound to a solid phase during amplification. The different oligonucleotide and/or PNA-oligomer sequences can be arranged on a plane  
10 solid phase in the form of a rectangular or hexagonal lattice, the solid phase surface preferably being composed of silicon, glass, polystyrene, aluminum, steel, iron, copper, nickel, silver, or gold, it being possible for other materials such as nitrocellulose or plastics to be used as well.

15 The fragments obtained by means of the amplification can carry a directly or indirectly detectable label. Preferred are labels in the form of fluorescence labels, radionuclides, or detachable molecule fragments having a  
20 typical mass which can be detected in a mass spectrometer, it being preferred that the fragments that are produced have a single positive or negative net charge for better detectability in the mass spectrometer. The detection may be carried out and visualized by means of matrix  
25 assisted laser desorption/ionization mass spectrometry (MALDI) or using electron spray mass spectrometry (ESI).

The amplicates obtained in the third step of the method are subsequently hybridized to an array or a set of oligonucleotides and/or PNA probes. In this context, the hybridization takes place in the manner described in the  
30 following. The set of probes used during the hybridization is preferably composed of at least 10 oligonucleotides or PNA-oligomers. In the process, the amplicates serve as probes which hybridize to oligonucleotides previously bonded to a solid phase. Ina particularly pre-  
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ferred embodiment, the oligonucleotides are taken from the group comprising Seq IDs 98 to 523. The non-hybridized fragments are subsequently removed. Said oligonucleotides contain at least one base sequence having a length of 10 nucleotides which is reverse complementary or identical to a segment of the base sequences specified in the appendix, the segment containing at least one CpG dinucleotide. The cytosine of the CpG dinucleotide is the 5<sup>th</sup> to 9<sup>th</sup> nucleotide from the 5'-end of the 10-mer. One oligonucleotide exists for each CpG dinucleotide.

In the next step of the method, the non-hybridized amplicates are removed.

In the final step of the method, the hybridized amplicates are detected. In this context, it is preferred that labels attached to the amplicates are identifiable at each position of the solid phase at which an oligonucleotide sequence is located.

According to the present invention, it is preferred that the labels of the amplicates are fluorescence labels, radionuclides, or detachable molecule fragments having a typical mass which can be detected in a mass spectrometer. The mass spectrometer is preferred for the detection of the amplicates, fragments of the amplicates or of probes which are complementary to the amplicates, it being possible for the detection to be carried out and visualized by means of matrix assisted laser desorption/ionization mass spectrometry (MALDI) or using electron spray mass spectrometry (ESI). The produced fragments may have a single positive or negative net charge for better detectability in the mass spectrometer.

The aforementioned method is preferably used for ascertaining genetic and/or epigenetic parameters of genomic DNA.

5 In order to enable this method, the invention further provides the chemically modified DNA of the genes estrogen receptor, p21, p27, p16, progesteron receptor, myoglobin, pcna, cdc2, c-erbB2, p53 and CEA as well as oligonucleotides and/or PNA-oligomers for detecting cytosine  
10 methylations. The present invention is based on the discovery that genetic and epigenetic parameters and, in particular, the cytosine methylation patterns of genomic DNA are particularly suitable for characterisation, grading, staging, and/or diagnosis of colon cancer.

15 The nucleic acids according to the present invention of Seq. ID No.12 through Seq. ID No. 523 can be used for characterisation, grading, staging and/or diagnosis of genetic and/or epigenetic parameters of genomic DNA.

20 This objective is achieved according to the present invention using a nucleic acid containing a sequence of at least 18 bases in length of the chemically pretreated genomic DNA according to one of Seq. ID No.32 through Seq.  
25 ID No.75 and sequences complementary thereto.

The chemically modified nucleic acid could heretofore not be connected with the ascertainment of disease relevant genetic and epigenetic parameters.

30 The object of the present invention is further achieved by an oligonucleotide or oligomer for the analysis of pretreated DNA, for detecting the genomic cytosine methylation state, said oligonucleotide containing at least  
35 one base sequence having a length of at least 10 nucleotides which hybridizes to a pretreated genomic DNA ac-

according to Seq. ID No.32 through Seq. ID No. 75. The oligomer probes according to the present invention constitute important and effective tools which, for the first time, make it possible to ascertain specific genetic and epigenetic parameters of colon cancers, in particular, for use in characterisation, grading, staging, and/or diagnosis of colon cancer. The base sequence of the oligomers preferably contains at least one CpG dinucleotide. The probes may also exist in the form of a PNA (peptide nucleic acid) which has particularly preferred pairing properties. Particularly preferred are oligonucleotides according to the present invention in which the cytosine of the CpG dinucleotide is the 5<sup>th</sup> - 9<sup>th</sup> nucleotide from the 5'-end of the 13-mer; in the case of PNA-oligomers, it is preferred for the cytosine of the CpG dinucleotide to be the 4<sup>th</sup> - 6<sup>th</sup> nucleotide from the 5'-end of the 9-mer.

The oligomers according to the present invention are normally used in so called "sets" which contain at least one oligomer for each of the CpG dinucleotides of the sequences of Seq. ID No. 32 to Seq. ID No. 75. Preferred is a set which contains at least one oligomer for each of the CpG dinucleotides from one of Seq. ID No. 32 to Seq. ID No. 75 .

In the case of the sets of oligonucleotides according to the present invention, it is preferred that at least one oligonucleotide is bound to a solid phase. It is further preferred that all the oligonucleotides of one set are bound to a solid phase.

The present invention moreover relates to a set of at least 10  $n$  (oligonucleotides and/or PNA-oligomers) used for detecting the cytosine methylation state in chemically pretreated genomic DNA (Seq. ID No.32 to Seq. ID No.75

No.75 and sequences complementary thereto). These probes enable characterisation, grading, staging and/or diagnosis of genetic and epigenetic parameters of colon cancer. Furthermore, the probes enable the diagnosis of predisposition to colon cancer. The set of oligomers may also be used for detecting single nucleotide polymorphisms (SNPs) in pretreated genomic DNA according to one of Seq. ID No. 32 to Seq. ID No. 75.

According to the present invention, it is preferred that an arrangement of different oligonucleotides and/or PNA-oligomers (a so-called "array") made available by the present invention is present in a manner that it is likewise bound to a solid phase. This array of different oligonucleotide- and/or PNA-oligomer sequences can be characterized in that it is arranged on the solid phase in the form of a rectangular or hexagonal lattice. The solid phase surface is preferably composed of silicon, glass, polystyrene, aluminum, steel, iron, copper, nickel, silver, or gold. However, nitrocellulose as well as plastics such as nylon which can exist in the form of pellets or also as resin matrices are possible as well.

Therefore, a further subject matter of the present invention is a method for manufacturing an array fixed to a carrier material for the grading, staging, and/or diagnosis of colon cancer, in which method at least one oligomer according to the present invention is coupled to a solid phase. Methods for manufacturing such arrays are known, for example, from US Patent 5,744,305 by means of solid-phase chemistry and photolabile protecting groups.

A further subject matter of the present invention relates to a DNA chip for the characterisation, grading, staging, and/or diagnosis of colon cancer. Furthermore the DNA chip enables the diagnosis of predisposition to colon



cancer. The DNA chip contains at least one nucleic acid according to the present invention. DNA chips are known, for example, in US Patent 5,837,832.

5 Moreover, a subject matter of the present invention is a kit which may be composed, for example, of a bisulfite-containing reagent, a set of primer oligonucleotides containing at least two oligonucleotides whose sequences in  
10 each case correspond or are complementary to a 18 base long segment of the base sequences specified in the appendix (Seq. ID No.32 through Seq. ID No.75), oligonucleotides and/or PNA-oligomers as well as instructions for carrying out and evaluating the described method. However, a kit along the lines of the present invention  
15 can also contain only part of the aforementioned components.

The oligomers according to the present invention or arrays thereof as well as a kit according to the present  
20 invention are intended to be used for the characterisation, grading, staging and/or diagnosis of colon cancer, or diagnosis of predisposition to colon cancer. According to the present invention, the method is preferably used for the analysis of important genetic and/or epigenetic  
25 parameters within genomic DNA, in particular for use in characterisation, grading, staging and/or diagnosis of colon cancer, and predisposition to colon cancer.

The methods according to the present invention are used,  
30 for example, for characterisation, grading, staging and/or diagnosis of colon cancer.

A further embodiment of the invention is a method for the analysis of the methylation status of genomic DNA without  
35 the need for chemical pretreatment. In the first step of the method the genomic DNA sample must be isolated from

tissue or cellular sources. Such sources may include cell lines, histological slides, body fluids, or tissue embedded in paraffin; for example, brain, central nervous system or lymphatic tissue. Extraction may be by means that are standard to one skilled in the art, these include the use of detergent lysates, sonification and vortexing with glass beads. Once the nucleic acids have been extracted the genomic double stranded DNA is used in the analysis.

In a preferred embodiment the DNA may be cleaved prior to the chemical treatment, this may be any means standard in the state of the art, in particular with restriction endonucleases. In the second step, the DNA is then digested with methylation sensitive restriction enzymes. The digestion is carried out such that hydrolysis of the DNA at the restriction site is informative of the methylation status of a specific CpG dinucleotide.

In the third step the restriction fragments are amplified. In a preferred embodiment this is carried out using a polymerase chain reaction.

In the final step the amplicates are detected. The detection may be by any means standard in the art, for example, but not limited to, gel electrophoresis analysis, hybridisation analysis, incorporation of detectable tags within the PCR products, DNA array analysis, MALDI or ESI analysis.

The present invention moreover relates to the diagnosis and/or prognosis of events which are disadvantageous or relevant to patients or individuals in which important genetic and/or epigenetic parameters within genomic DNA, said parameters obtained by means of the present invention may be compared to another set of genetic and/or epigenetic parameters, the differences serving as the ba-

sis for a diagnosis and/or prognosis of events which are disadvantageous or relevant to patients or individuals.

5 In the context of the present invention the term "hybridization" is to be understood as a bond of an oligonucleotide to a completely complementary sequence along the lines of the Watson-Crick base pairings in the sample DNA, forming a duplex structure.

10 The term "functional variants" denotes all DNA sequences which are complementary to a DNA sequence, and which hybridize to the reference sequence under stringent conditions.

15 In the context of the present invention, "genetic parameters" are mutations and polymorphisms of genomic DNA and sequences further required for their regulation. To be designated as mutations are, in particular, insertions, deletions, point mutations, inversions and polymorphisms  
20 and, particularly preferred, SNPs (single nucleotide polymorphisms).

In the context of the present invention, "epigenetic parameters" are, in particular, cytosine methylations and  
25 further chemical modifications of DNA bases of genomic DNA and sequences further required for their regulation. Further epigenetic parameters include, for example, the acetylation of histones which, cannot be directly analyzed using the described method but which, in turn, correlates with the DNA methylation.  
30

In the following, the present invention will be explained in greater detail on the basis of the sequences and examples without being limited thereto.

Seq. ID 1 to 11 represent the genomic DNA of genes estrogen receptor, p21, p27, p16, progesteron receptor, myoglobin, pcna, cdc2, c-erbB2, p53 and CEA. These sequences are derived from Genbank and will be taken to include all minor variations of the sequence material which are currently unforeseen, for example, but not limited to, minor deletions and SNPs.

Sequence ID 12 to 31 represent segments of genomic DNA which are particularly useful for the determination of colon cell proliferative disorder.

Sequence ID 32 to 75 exhibit the chemically pretreated sequence of genes estrogen receptor, p21, p27, p16, progesteron receptor, myoglobin, pcna, cdc2, c-erbB2, p53 and CEA. These sequences will be taken to include all minor variations of the sequence material which are currently unforeseen, for example, but not limited to, minor deletions and SNPs.

Sequences having even sequence numbers (e.g., Seq. ID No. 32, 34, 36, ...) exhibit in each case sequences of chemically pretreated genomic DNAs.

Sequences having odd sequence numbers (e.g., Seq. ID No. 33, 35, 37 ...) exhibit in each case the sequences of chemically pretreated genomic DNAs. Said genomic DNAs are complementary to the genomic DNAs from which the preceding sequence was derived (e.g., the complementary sequence to the genomic DNA from which Seq. ID No.32 is derived is the genomic sequence from which Seq. ID No.33 is derived, the complementary sequence to the genomic DNA from which Seq. ID No.33 is derived is the sequence from which Seq. ID No.34 is derived, etc.)

Sequence ID 76 to 97 exhibit the sequence of primer oligonucleotides for the amplification of chemically pretreated DNA according to Sequence IDs 32 to 75.

- 5     Sequence IDs 98 to 523 exhibit the sequence of oligomers which are particularly useful for the analysis of CpG positions within chemically pretreated DNA according to Sequence IDs 32 to 75.
- 10    The following examples describe the invention in detail without limiting the scope of the invention.

Example 1: Description of PCR

- 15    The single gene PCR reaction was performed using a thermocycler (Eppendorf GmbH) using 10 ng of bisulfite treated DNA, 6 pmole of each primer, 200 µM of each dNTP, 1.5 mM MgCl<sub>2</sub> and 1 U of HotstartTaq (Qiagen AG). The other conditions were as recommended by the Taq polymerase manufacturer. Single genes were amplified by PCR
- 20    performing a first denaturation step for 14 min at 96 °C, followed by 39 cycles (60 sec at 96°C, 45 sec at 55 °C, 75 sec at 72 °C) and a subsequent final elongation of 10 min at 72 °C. The bisulfite DNA was prepared according to a published procedure from genomic DNA individually isolated from 12 matched samples of adenocarcinoma of the
- 25    colon and healthy colon tissue. The genomic DNA was isolated using the wizzard DNA isolation kit (Promega, Madison).

- 30    Example 2 : Methylation analysis of gene p16.

The following example relates to a fragment of the gene p16 in which a specific CG dinucleotide is to be analyzed for methylation.

- 35    In the first step, a genomic sequence is treated using bisulfite (hydrogen sulfite, disulfite) in such a manner

that all cytosines which are not methylated at the 5-position of the base are modified in such a manner that a different base is substituted with regard to the base pairing behavior while the cytosines methylated at the 5-position remain unchanged.

If bisulfite solution is used for the reaction, then an addition takes place at the non-methylated cytosine bases. Moreover, a denaturing reagent or solvent as well as a radical interceptor must be present. A subsequent alkaline hydrolysis then gives rise to the conversion of non-methylated cytosine nucleobases to uracil. The chemically converted DNA is then used for the detection of methylated cytosines. In the second method step, the treated DNA sample is diluted with water or an aqueous solution. Preferably, the DNA is subsequently desulfonated. In the third step of the method, the DNA sample is amplified in a polymerase chain reaction, preferably using a heat-resistant DNA polymerase. In the present case, cytosines of the gene p16 are analyzed. To this end, a defined fragment having a length of 598 bp is amplified with the specific primer oligonucleotides TTGAAAATTAAGGGTTGAGG (Sequence ID 82) and CACCCTCTAATAACCAACCA (Sequence ID No. 83).

The amplificate serves as a sample which hybridizes to an oligonucleotide previously bound to a solid phase, forming a duplex structure, for example TAAGTGTTCGGAGTTAAT (SEQ ID NO: 238), the cytosine to be detected being located at position 439 of the amplificate. The detection of the hybridization product is based on Cy3 and Cy5 fluorescently labelled primer oligonucleotides which have been used for the amplification. A hybridization reaction of the amplified DNA with the oligonucleotide takes place only if a methylated cytosine was present at this location in the bisulfite-treated DNA as shown for healthy

tissue in Figure 1A. Thus, the methylation status of the specific cytosine to be analyzed is inferred from the hybridization product.

5 In order to verify the methylation status of the position, a sample of the amplificate is further hybridized to another oligonucleotide previously bonded to a solid phase. Said oligonucleotide is identical to the oligonucleotide previously used to analyze the methylation  
10 status of the sample, with the exception of the position in question. At the position to be analysed said oligonucleotide comprises a thymine base as opposed to a cytosine base i.e TAAGTGTTTGGAGTTAAT (SEQ ID NO: 239). Therefore, the hybridisation reaction only takes place if an  
15 unmethylated cytosine was present at the position to be analysed as shown for tumor tissue in Figure 1B.

Example 3: Differentiation between colon tumour and healthy colon tissue

20 Differentiation of healthy samples and adenocarcinoma tumours. For tumour class prediction between healthy and tumor tissue we used a Support Vector Machine (SVM) on a set of selected CpG sites (F. Model, P. Adorjan, A. Olek, C. Piepenbrock, Feature selection for DNA methylation based  
25 cancer classification. Bioinformatics. 2001 Jun;17 Suppl 1:S157-64.). First we ranked the CpG sites for a given separation task by their significance of the difference between the two class means. The significance of each CpG was estimated by a two sample t-test (W, Mendenhall, T,  
30 Sincich, Statistics for engineering and the sciences (Prentice-Hall, New Jersey 1995).

In order to relate the methylation patterns to a adenocarcinoma tumour, it is initially required to comparatively analyze the DNA methylation patterns of healthy  
35 tissue and adenocarcinoma tumours tissue (Figure 2 A and

B). These analyses were carried out, analogously to Examples 1. The results obtained in this manner are stored in a database and the CpG dinucleotides which are methylated differently between the two groups are identified. This  
5 can be carried out by determining individual CpG methylation rates as can be done, for example, by sequencing, which is a relatively imprecise method of quantifying methylation at a specific CpG, or else, in a very precise manner, by a methylation-sensitive "primer extension reaction". In a particularly preferred variant, as illustrated in the preceeding examples the methylation status of hundreds or thousands of CpGs may be analysed on an oligomer array. It is also possible for the patterns to be compared, for example, by clustering analyses which  
10 can be carried out, for example, by a computer.

A panel of genomic fragments of 11 different genes (listed in Table 1) were bisulphite treated and amplified by singleplex PCRs according to Example 1. However, as  
20 will be obvious to one skilled in the art, it is also possible to use other primers that amplify the genomic, bisulphite treated DNA in an adequate manner, and/or to carry out the PCRs in a multiplex format. However the primer oligonucleotide pairs as listed in Table 1 are  
25 particularly preferred. In order to differentiate adenocarcinoma tumour from healthy control samples optimal results were obtained by including at least 6 CpG dinucleotides, the most informative CpG positions for this discrimination being located within the p16, p53, CEA, c-erbB2 and estrogen receptor genes (cf. Fig. 2, Tab1). In  
30 addition, the majority of the analysed CpG dinucleotides of the panel showed different methylation patterns between the two phenotypes. The results prove that methylation fingerprints are capable of providing differential  
35 diagnosis of adenocarcinoma tumours and could therefore be applied in a large number clinical situations



For class prediction a SVM was trained on the most significant CpG positions, where the optimal number of CpG sites depends on the complexity of the separation task. Implementation of the SVM used the Sequential Minimal Optimization algorithm to find the 1-norm soft margin separating hyperplane (N. Christianini, J. Shawe-Taylor, An Introduction to Support Vector Machines, Cambridge University Press, Cambridge 2000). The box constraint was set to  $C=10$ . Generalization performance was estimated by averaging over 50 cross validation runs on randomly permuted samples partitioned into 8 groups.

Example 4: Analysis of the methylation status of the most informative CpG positions of the genes c-erbB2, p53, CEA, p16 and ER1

The methylation status of the most informative CpG positions of the gene fragments of genes c-erbB2, p53, CEA, p16 and ER1 are shown in this example. Corresponding to Example 2, where the methylation status is demonstrated by spots, Table 2 describes in a more detailed way the methylation status of different gene fragments of various patients by calculating the methylation status of colon tumour and healthy colon tissue. The first column indicates the specific gene fragment, the second column describes the investigated CpG Oligonukleotide, the third column depicts the diagnosis of the investigated tissue (T=tumor, H=healthy) and columns 4 to 17 show the logarithm of the ratio of the fluorescence signal of the CG oligonucleotide versus TG oligonucleotide of colon tumour and healthy colon tissue of 14 different patients. For example, a comparison of the methylation status of gene p16, patient 11, shows that the healthy tissue is less methylated compared to the tumour tissue for this sample. The opposite ratio can be observed, for example, for gene c-erbB2 for patient 11. In this case the tumour sample is

more methylated than the healthy sample. The analyzed CpG positions show that the genes p53, CEA, p16 and ER1 are hypermethylated, whereas c-erbB2 is hypomethylated in most of the tumour samples compared with the healthy controls.

Example 5: Identification of the methylation status of CpG sites of genes CEA and p16 by methylation sensitive restriction enzyme digest.

In the CEA gene, a defined fragment having a length of 351 bp, which contains 7 CpG sites, is amplified with the specific primer oligonucleotides TGGTTAAATGTGTGGGAGAT (Sequence ID 524) and TCCTGAGTGATGTCTGTGTG (Sequence ID No. 525) and in the p16 gene, a defined fragment having a length of 391 bp, which contains 26 CpG sites, is amplified with the specific primer oligonucleotides ATGACACCAAACACCCCGAT (Sequence ID 526) and CTGTCCCTCAAATCCTCTG (Sequence ID No. 527). CGCG for gene CEA with Cytosins at positions 127 and 129 of the amplificate and CGCG for gene p16 with Cytosins at positions 362 and 364 of the amplificate, are located in a SacII restriction enzyme recognition sequence, CCGCGG. The cleavage of SacII is blocked by methylation of at least one of the two CpG dinucleotides.

The genomic DNA isolated from adenocarcinoma of colon tissue and from healthy colon tissue was hydrolysed by SacII as recommended by the manufacturer (New England Biolabs GmbH).

10 ng of the SacII restricted DNA was used as template for the amplification of the above indicated CEA and p16 gene fragments. The PCR reaction was performed using a thermocycler (Eppendorf GmbH) using 10 ng of DNA, 6 pmole of each primer, 200 µM of each dNTP, 1.5 mM MgCl<sub>2</sub> and 1 U

of HotstartTaq (Qiagen AG). The other conditions were as recommended by the Taq polymerase manufacturer. Using the above mentioned primers, gene fragments were amplified by PCR performing a first denaturation step for 14 min at 96 °C, followed by 30 - 45 cycles (step 2: 60 sec at 96°C, step 3: 45 sec at 55 °C , step 4: 75 sec at 72 °C) and a subsequent final elongation of 10 min at 72 °C. The presence of PCR products was analysed by agarose gel electrophoresis.

PCR products were detectable with SacII hydrolyzed DNA isolated from colon cancer tissue, when step 2 to step 4 of the cycle program were repeated 34, 37, 39, 42 and 45 fold. In contrast PCR products were only detectable with SacII hydrolyzed DNA isolated from healthy colon tissue when step 2 to step 4 of the cycle program were repeated 42 and 45 fold. These results indicate that at least one of CpG positions located within the SacII recognition sequence of the analysed CEA and the p16 gene fragment showed a higher methylation status in cancer samples compared to the healthy control.

#### Description of figures

##### Figure 1

Figure 1 shows the hybridisation of fluorescent labelled amplicates to a surface bound oligonucleotide. Sample A being from healthy tissue and sample B being from colon adenocarcinoma tissue. Fluorescence at a spot, denoted by an arrow, indicates hybridisation of the amplicate against the oligonucleotide. Hybridisation to a CG oligonucleotide with the sequence TAAGTGTTCGGAGTTAAT (SEQ ID NO: 238) denotes methylation at the cytosine position being analysed, hybridisation to a TG oligonucleotide with the sequence TAAGTGTTGGAGTTAAT (SEQ ID NO: 239) denotes no methylation at the cytosine position be-

ing analysed. It can be seen that sample A was umethy-  
lated for CG positions of the amplificate of gene p16  
whereas in comparison sample B had a higher degree of me-  
thylation at the same position.

5

#### Figure 2

Differentiation of colon tumour(A) from healthy colon  
tissue(B). High probability of methylation corresponds to  
red, uncertainty to black and low probability to green.

10

The labels on the left side of the plot are gene (e.g.  
for the topmost: 2064) and CpG (e.g. for the topmost:  
1485A) identifiers. The hybridisation was carreid out  
with Cy5 labelled amplificates generated by singlplex PCR  
reactions using primer oligonucleotides as shown in Table

15

1. The labels on the right side give the significance (p-  
value, T-test) of the difference between the means of the  
two groups. Each row corresponds to a single CpG and each  
column to the methylation levels of one sample. CpGs are  
ordered according to their contribution to the distinc-  
tion to the differential diagnosis of the two lesions  
with increasing contribution from top to bottom.

20

Table 1  
List of genes, reference numbers (ID) according Fig 2 and primer oligonucleotides according to Example 2 and Figures 1 and 2.

Gene	Gene Seq-ID	Accession no	Primer Seq-ID	PCR primer	Primer Seq-ID	PCR primer
Estrogen-receptor	1	NM_000125	76	AGGAGGGGGAATTAAATAGA	77	ACAATAAAACCATCCCAAAATAC
	2	NM_000389	78	GGATTAGTGGGAATAGAGGTG	79	AAACCCAAACTCTCTAACTACC
	3	NM_004084	80	GTGGGAGGTAGTTGAAGA	81	ATACACCCCTAACCCCAAAAT
	4	NM_000077	82	TTGAAAATTAAAGGGTTGAGG	83	CACCTCTAATAACCAACCA
Progesteron-receptor	5	NM_000926	84	GAGGGGTAGTGGAAATTAG	85	CCTTACCTTCAACTCAATCA
Myoglobin	6	NM_005368	86	GTTTTGGTAAAGGGGTAGAA	87	CCTAAAAATATCAACCTCCACCT
PCNA	7	NM_002592	88	TTTTAGGTTGTAAAGGAGGTTT	89	TAAATACCTCCCAACACCTTTCT
CDC2	8	NM_001786	90	ATTAGAAGTGAAGTAATGGAATTT	91	TCAATTTCCAAAAACCAAC
c-erbB2	9	NM_004448	92	GGAGGGGTAGAGTTATTAGTT	93	TATACCTCTCAAAACCAACCTC
P53	10	NM_000546	94	GATTGGTAAGTTTTTGATTGA	95	AAATCTCCCAACAATACAACTC
CEA	11	NM_004363	96	GTTAGGATGGGATTAAAGTGTG	97	AATCAAAATATCCCAAAATACAA

Table 2

gene	QpG	Diagnosis	log (fluorescence OGoligo / fluorescence TGoligo) of matched pair of odontotumor (T) and healthy odontotissue (H) <sup>a</sup>															
			11	3	9	1	8	2	4	13	12	14	15	5	10	6		
c-erbB2	2064:148	T	-1,07	-1,72	-1,11	-1,53	-1	-1,3	-1,63	-1,01	-1,64	-1,22	-1,33	-0,88	-1,57	-1,22		
		H	-0,82	-1,09	-1,34	-1,17	-0,75	-1,09	-1,36	-0,89	-0,88	-0,72	-1,06	-0,93	-1,43	-0,7		
p53	2317:122	T	-2,17	-0,58	-2,37	-1,91	-2,08	-0,32	-0,3	-1,63	-2,19	-1,87	-1,71	-4,31	-3,15	-1		
		H	-4,03	-3,01	-3,14	-3,83	-1,53	-2,29	-1,86	-2,96	-2,76	-4,09	-3,44	-4,77	-2,33	-3,97		
p53	2317:153	T	-2,38	-2,84	-2,77	-2,57	-2,93	-2,44	-2,89	-3,12	-2,77	-2,32	-2,7	-2,89	-2,13	-2,29		
		H	-3,36	-3,36	-3,17	-3,32	-3,15	-3,47	-2	-2,94	-3,8	-3,77	-3,67	-4,12	-2,2	-3,12		
CEA	2398:176	T	-2,76	-1,84	-3,7	-2,42	-2,59	-0,83	-2,14	-1,96	-2,86	-4,02	-2,76	-5,71	-5,07	-2,33		
		H	-4,32	-2,64	-4,72	-3,9	-4,43	-4,85	-2,95	-3	-2,67	-4,19	-2,92	-4,63	-4,53	-3,64		
CEA	2398:227	T	-3,35	-2,15	-3,83	-3,88	-4,02	-2,95	-2,81	-3,98	-3,9	-4,01	-4,75	-4,64	-4,34	-3,15		
		H	-4,7	-4,37	-5,1	-5,77	-4,7	-4,64	-3,33	-3,65	-5,48	-5,2	-5,32	-5,75	-3,84	-5,4		
p16	2035:181	T	-1,64	-1,99	-2,66	-2,6	-3,78	-1,07	-2,21	-2,18	-3,24	-1,8	-2,19	-3,21	-3,25	-2,13		
		H	-2,74	-3,02	-4,09	-3,52	-3,74	-3,87	-2,88	-2,76	-3,28	-2,27	-3,19	-3,38	-3,88	-3,49		
ER1	41:2912	T	-0,4	-0,37	-1,23	-0,96	-1,36	-0,47	0,37	-0,34	-0,76	-0,85	-0,56	-1,32	-1,53	-1,33		
		H	-0,8	-1,25	-2,1	-1,23	-1,52	-1,38	-0,44	-0,93	-1,26	-1,55	-1,21	-1,55	-1	-1,45		
ER1	41:2860	T	-1,06	-0,77	-2,06	-1,8	-1,7	-0,53	-1,52	-1,82	-1,19	-1,59	-0,82	-1,6	-1,85	-1,69		
		H	-1,83	-2,03	-2,05	-2,22	-2,36	-2,13	-2,26	-1,72	-0,91	-2,11	-1,6	-2,05	-1,9	-2,14		
ER1	41:2428	T	0,02	0,84	-1,41	-1,22	-1,39	0,67	-0,13	-0,33	-0,66	-1,41	0,22	-1,78	-1,61	-1		
		H	-0,97	-0,86	-1,61	-1,36	-1,02	-1,78	-0,88	-1,05	-1,65	-1,29	-1,19	-1,53	-1,45	-1,96		
ER1	41:2849	T	-0,86	-0,43	-1,55	-0,98	-1,22	-0,97	-2,16	-1,2	-0,66	-1,07	-0,54	-1,59	-1,45	-0,78		
		H	-1,11	-1,04	-1,97	-1,27	-2,06	-2,21	-2,77	-1,01	-1,08	-1,59	-1,08	-1,9	-2,02	-1,76		

a) the values indicate the mean of at least 12 O3TGoligo pairs analysed in 3 independent chip hybridisation

1. A method to determine the methylation status of CpG dinucleotides within one or more of the genes estrogen receptor, p21, p27, p16, progesterone receptor, myoglobin, pcna, cdc2, c-erbB2, p53 and CEA comprising contacting the target nucleic acid in a biological sample with at least one reagent or series of reagents wherein said reagent or series of reagents distinguishes between methylated and non methylated CpG dinucleotides within the target nucleic acid and concluding from the methylation status of one or more of said CpG positions on the presence or absence of a colon cell proliferative disorder.
2. A method according to Claim 1 comprising the following steps:
  - obtaining a biological sample containing genomic DNA
  - extracting the genomic DNA
  - in the genomic DNA sample, cytosine bases which are unmethylated at the 5-position are converted, by treatment, to uracil or another base which is dissimilar to cytosine in terms of base pairing behavior;
  - fragments of the pretreated genomic DNA are amplified using sets of primer oligonucleotides according to Seq ID 76 to Seq ID 97 and a polymerase, the amplicates carrying a detectable label;
  - detection of the fragments
  - Identification of the methylation status of one or more cytosine positions
3. A method according to claim 2, characterized in that the reagent is a solution of bisulfite, hydrogen sulfite or disulfite.

4. A method as recited in Claims 2 and 3,  
characterized in that the amplification is carried  
out by means of the polymerase chain reaction (PCR).
- 5 5. A method as recited in one of the Claims 2 to 4,  
characterized in that more than ten different frag-  
ments having a length of 100 - 2000 base pairs are  
amplified.
- 10 6. A method as recited in one of the Claims 2 to 5,  
characterized in that the amplification of several  
DNA segments is carried out in one reaction vessel.
- 15 7. A method as recited in one of the Claims 2 to 6,  
characterized in that the polymerase is a heat-  
resistant DNA polymerase.
- 20 8. A method as recited in one of the Claims 2 to 7,  
characterized in that the labels of the amplicates  
are fluorescence labels.
- 25 9. A method as recited in one of Claims 2 to 7,  
characterized in that the labels of the amplicates  
are radionuclides.
- 30 10. A method according to one of Claims 2 to 9, charac-  
terized in that each amplicate is detected by hy-  
bridization to an oligonucleotide or peptide nucleic  
acid (PNA)-oligomer.
- 35 11. A method according to claim 10, characterized in that  
the oligonucleotide or peptide nucleic acid (PNA)-  
oligomer is taken from the group comprising Seq ID 98  
to 523.



12. A method as recited in one of Claims 2 to 7,  
characterized in that the labels of the amplificates  
are detachable molecule fragments having a typical  
mass which are detected in a mass spectrometer.
13. A method as recited in one of Claims 2 to 7 and 12,  
characterized in that the amplificates or fragments  
of the amplificates are detected in the mass spec-  
trometer.
14. A method as recited in one of Claims 12 or 13, char-  
acterized in that the produced fragments have a sin-  
gle positive or negative net charge for better de-  
tectability in the mass spectrometer.
15. A method as recited in one of the Claims 12 through  
14, characterized in that detection is carried out  
and visualized by means of matrix assisted laser de-  
sorption/ionization mass spectrometry (MALDI) or us-  
ing electron spray mass spectrometry (ESI).
16. A method as recited in Claim 2, characterized in that  
the amplification step preferentially amplifies DNA  
which is of particular interest in healthy and/or  
diseased colon tissues, based on the specific genomic  
methylation status of colon tissue, as opposed to  
background DNA.
17. A method according to Claim 1 comprising the follow-  
ing steps;  
a) obtaining a biological sample containing genomic  
DNA  
b) extracting the genomic DNA,  
c) digesting the target nucleic acids with one or  
more methylation sensitive restriction enzymes,  
d) amplification of the DNA digest and

e) detection of the amplificates.

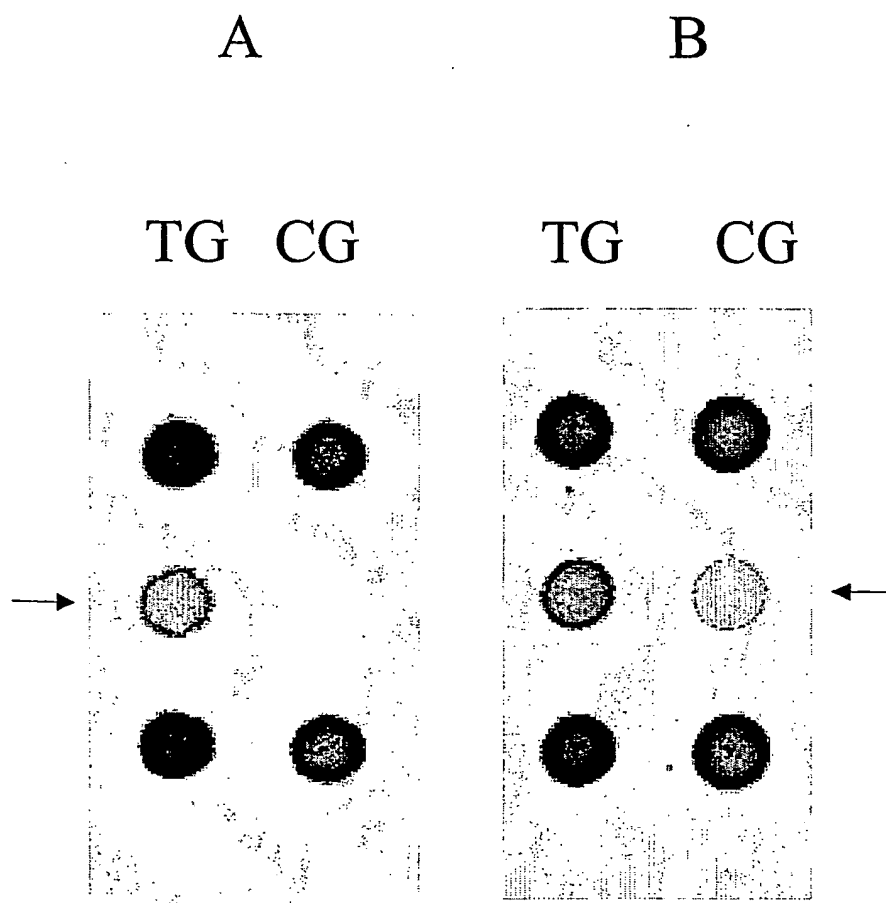
- 5 18. A method according to Claim 17 wherein the target nucleic acids comprise one or more sequences taken from the group according to Seq ID 12 to Seq ID 31 or sequences hybridising thereto and fragments thereof.
- 10 19. A method as recited in one of Claims 17 or 18, characterized in that the amplification is carried out by means of the polymerase chain reaction (PCR).
- 15 20. A method as recited in one of Claims 17 to 19, characterized in that the amplification of several DNA segments is carried out in one reaction vessel.
21. A method as recited in one of Claims 17 to 20, characterized in that the polymerase is a heat-resistant DNA polymerase.
- 20 22. An isolated nucleic acid of the pretreated genomic DNA according to one of the sequences taken from the group comprising Seq. ID No. 32 to Seq. ID No. 75 and sequences complementary thereto.
- 25 23. An oligomer, in particular an oligonucleotide or peptide nucleic acid (PNA)-oligomer, said oligomer comprising at least one base sequence of at least 10 nucleotides which hybridizes to or is identical to a pretreated genomic DNA according to one of the Seq. ID No. 32 to Seq. ID No 75 according to claim 22.
- 30 24. An oligomer or peptide nucleic acid (PNA)-oligomer as recited in Claim 23, wherein the base sequence includes at least one CpG dinucleotide sequence.
- 35

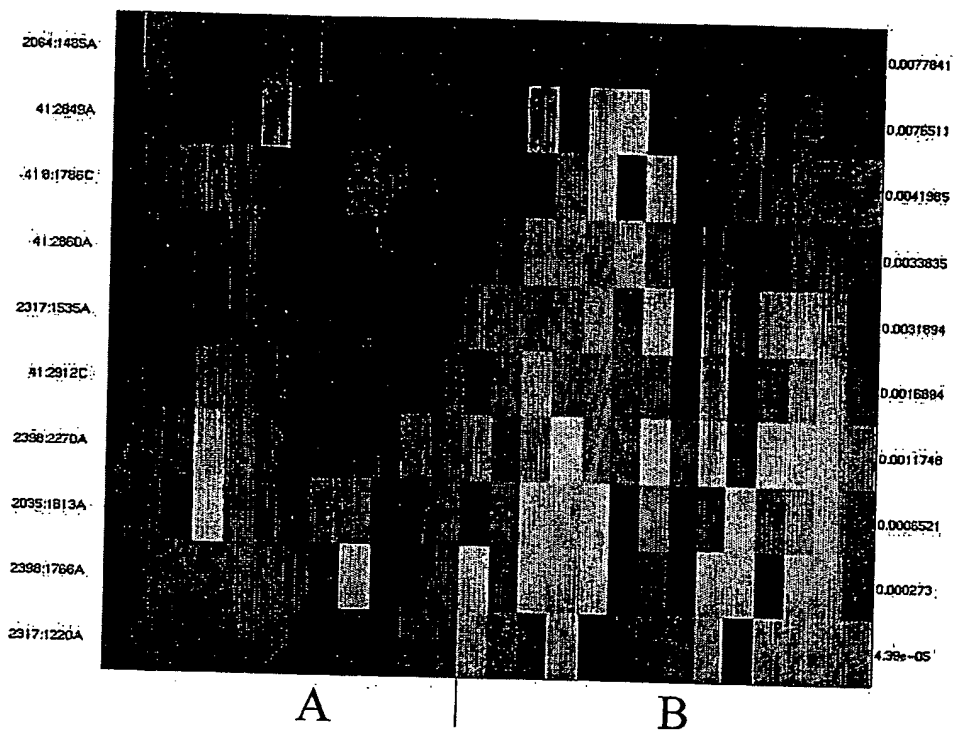
- 5 25. An oligomer or peptide nucleic acid (PNA)-oligomer as recited in Claim 23, characterized in that the cytosine of the at least one CpG dinucleotide is/are located approximately in the middle third of the oligomer.
- 10 26. An oligomer or peptide nucleic acid (PNA)-oligomer, in particular an oligonucleotide, according to one of the sequences taken from the group comprising Seq. ID No.98 to Seq. ID No. 523.
- 15 27. A set of oligomers or peptide nucleic acid (PNA)-oligomers, comprising at least two oligomers according to any of claims 22 to 26.
- 20 28. A set of oligomers or peptide nucleic acid (PNA)-oligomers as recited in Claim 27, comprising oligomers for detecting the corresponding genomic methylation state of all CpG dinucleotides within one of the sequences according to Seq. ID Nos. 32 to 75 according to claim 22, and sequences complementary thereto.
- 25 29. A set of at least two oligonucleotides or peptide nucleic acid (PNA)-oligomers as recited in Claim 23, as primer oligonucleotides for the amplification of DNA sequences of one of Seq. ID 32 to Seq. ID 75 and/or sequences complementary thereto and segments thereof.
- 30 30. A set of oligonucleotides or peptide nucleic acid (PNA)-oligomers as recited in Claims 22 and 23, characterized in that at least one oligonucleotide is bound to a solid phase.
- 35 31. Use of a set of oligomers or peptide nucleic acid (PNA)-oligomers according to any of claims 22 to 25

as probes for determining the cytosine methylation state and/or single nucleotide polymorphisms (SNPs) of a corresponding genomic DNA by analysis of a chemically pretreated genomic DNA according to claim 2.

32. Use of a pretreated genomic DNA according to claim 22 for the determination of the methylation status of a corresponding genomic DNA and/or detection of single nucleotide polymorphisms (SNPs).
33. A method for manufacturing an arrangement of different oligomers or peptide nucleic acid (PNA)-oligomers (array) for analyzing diseases associated with the corresponding genomic methylation status of the CpG dinucleotides within one of the Seq. ID 32 to Seq. ID 75 and sequences complementary thereto, wherein at least one oligomer according to any of the claims 22 to 26 is coupled to a solid phase.
34. An arrangement of different oligomers or peptide nucleic acid (PNA)-oligomers (array) obtainable according to claim 33.
35. An array of different oligonucleotide- and/or PNA-oligomer sequences as recited in Claim 34, characterized in that these are arranged on a plane solid phase in the form of a rectangular or hexagonal lattice.
36. A DNA/PNA array for the analysis of prostate cell proliferative disorders associated with the methylation state of genes comprising at least one nucleic acid according to one of the preceding claims.

- 5 37. An array as recited in any of the Claims 34 to 36,  
characterized in that the solid phase surface is com-  
posed of silicon, glass, polystyrene, aluminium,  
steel, iron, copper, nickel, silver, or gold.
- 10 38. Use of a method according to one of Claims 1 through  
21 for the characterisation, classification, diagno-  
sis and differentiation of colon cell proliferative  
disorders.
- 15 39. A kit comprising a bisulfite (= disulfite, hydrogen  
sulfite) reagent as well as oligonucleotides and/or  
PNA-oligomers according to one of the Claims 22 to  
29.
- 20 40. Use of a pretreated genomic DNA according to claim  
22 for the characterisation, classification, diagno-  
sis and differentiation of colon cell proliferative  
disorders.

*Fig 1*

*Fig 2*

## Sequence listing

&lt;110&gt; Epigenomics AG

&lt;120&gt; Method and nucleic acids for the analysis of colon cancer

&lt;160&gt; 527

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&lt;212&gt; DNA

&lt;213&gt; Homo Sapiens

&lt;400&gt; 1

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&lt;213&gt; Homo Sapiens

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&lt;212&gt; DNA

&lt;213&gt; Homo Sapiens

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&lt;213&gt; Homo Sapiens

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&lt;211&gt; 3664

&lt;212&gt; DNA

&lt;213&gt; Homo Sapiens

&lt;400&gt; 8

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&lt;211&gt; 1729

&lt;212&gt; DNA

&lt;213&gt; Homo Sapiens

&lt;400&gt; 9

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&lt;213&gt; Homo Sapiens

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&lt;210&gt; 12

&lt;211&gt; 18

&lt;212&gt; DNA

&lt;213&gt; Homo Sapiens

&lt;400&gt; 12

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18



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&lt;210&gt; 27

&lt;211&gt; 18

&lt;212&gt; DNA

&lt;213&gt; Homo Sapiens

&lt;400&gt; 27

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18

&lt;210&gt; 28

&lt;211&gt; 18

&lt;212&gt; DNA

&lt;213&gt; Homo Sapiens

&lt;400&gt; 28

aagtgcccgcc ggctgccc

18

&lt;210&gt; 29

&lt;211&gt; 18

&lt;212&gt; DNA

&lt;213&gt; Homo Sapiens

&lt;400&gt; 29

gggcagccgc gggcactt

18

&lt;210&gt; 30

&lt;211&gt; 18

&lt;212&gt; DNA

&lt;213&gt; Homo Sapiens

&lt;400&gt; 30

ccctgagccg cagctcag

18

&lt;210&gt; 31

&lt;211&gt; 18

&lt;212&gt; DNA

&lt;213&gt; Homo Sapiens

&lt;400&gt; 31

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18

&lt;210&gt; 32

&lt;211&gt; 3315

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 32

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21

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&lt;210&gt; 33

&lt;211&gt; 3315

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 33

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&lt;210&gt; 34

&lt;211&gt; 7369

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 34

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ag	cgaggg	at	tttag	tgaa	tattag	480
tgaat	tttag	ttggag	gaaat	gagag	ggtgg	540
ggaag	gttgt	gcggt	ttagg	gaaat	taatt	600
aaaga	aaagt	gggac	tttat	gtta	atggt	660
ttgt	gaagga	tttgg	gggtg	tcgt	aaatta	720
tg	agttata	ttatt	gattg	attag	tttgt	780
tttt	ttgag	tttt	tattg	gag	ttatg	840
tg	tcgc	at	agaa	tcgt	tttt	900
gg	tttt	atggt	tttt	aggt	gaatt	960
ttatt	ttatt	tagt	acgt	gggg	ag	1020
gtaa	cgcg	ttcgt	acgag	ggc	gaggg	1080
ttggg	ttcgt	cgg	cgagt	gcg	cgg	1140
gttt	cgg	ggc	gggag	tcg	ggc	1200
cgaat	aacgt	cggg	at	tcg	tttc	1260
tttg	ttgt	taatt	tttt	ttag	taggt	1320
ggtag	tttt	agtt	ttag	tttc	gttc	1380
gcgc	cggac	gggag	ggaat	cggg	ttag	1440
gttc	cggt	cgg	cg	aatg	tttt	1500
taaat	cgtc	attat	tttc	tttt	atcgt	1560
tg	gggt	gttc	tcggg	tttt	tttt	1620
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cggg	ggg	tcg	tcg	ttg	tttc	1740
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gcg	cgg	tcg	tcg	tag	gtc	1860
gtc	ggg	gttc	tag	gttt	ggg	1920
ttgc	cggt	ttgt	ttgt	gag	gag	1980
cgtg	cggg	atgt	tcg	taac	tttag	2040
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gttc	cggt	ttac	at	gtat	gata	2160
agg	cgat	gcg	aat	taa	gagg	2220
acg	aggt	agag	aagg	tg	ttat	2280
tttt	taag	tg	agg	ggat	ggg	2340
cgg	cggt	ttta	at	tac	gtg	2400
agatt	gtc	tag	tag	atg	ata	2460
gatt	cgac	gatt	ta	atg	ggt	2520
tg	gggt	tttag	tttc	tatt	tag	2580
agag	ttatt	taggt	gtg	gtt	tg	2640
tg	att	ggag	agat	at	ttta	2700

28

cgcggttttt	tttattagta	attttttaggt	atgtgataaa	gttgggatgt	ttattaacgg	2760
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aggtagtggg	ttttttat	tttgattatg	gggttaattt	ttgttagtta	ttgttttttt	2940
taataaagat	tgtgtgtttt	ttttaaaaaa	tttttttgcg	tttaga		2986

&lt;210&gt; 37

&lt;211&gt; 2986

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 37

tttaagcgta	ggggaaat	ttaaaaagaa	tatataat	ttattagaaa	aaataatggt	60
tggtagaagt	tggttttata	gttaggggat	gaaaaat	ttatttttat	ttataat	120
ggattcggtta	tttttagtgg	tttttaagg	tataaaatga	ggttggtttt	aaaaaataga	180
aagtgtat	ttcgttttag	agttttttt	ttagtttagga	ggcggatcgt	tgataaatat	240
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gcgtattggt	tcgttaatt	cggttggtg	ttcgacggat	tagtttttg	gtttattaaa	600
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29

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atattatatt	atacgtttta	gaatgtgata	gttgaagg	attttt		2986

&lt;210&gt; 38

&lt;211&gt; 5666

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 38

aaaattagaa	tttttatttt	tttgcgtttg	ttatatTTTT	tagtggttgt	taattttttt	60
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&lt;210&gt; 39

&lt;211&gt; 5666

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 39

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cgtaaggagg	taaaagtttt	aatttt				5666

&lt;210&gt; 40

&lt;211&gt; 5085

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 40

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&lt;210&gt; 41

&lt;211&gt; 5085

&lt;212&gt; DNA



&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 41

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35

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&lt;210&gt; 42

&lt;211&gt; 8222

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 42

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&lt;210&gt; 43

&lt;211&gt; 8222

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 43

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g g t t t g a g t a	a a g a a g g a a	a a a t a g g g g t	g t g g a g t t t a	t t t a g t a g t t	t t t t t t t t t t	3900
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a a t t a g t t t a	g a a t g g t g a g	g g a a g g g g t t	a a g a g a a g a g	t g t a t t a t t t	t t g g g a t a g a	5160
a g g t a a c g a g	g a g a t g a g t t	t t t a t g t a t a	a a t t a t a g t t	a t t t t t t a t t	t a t a a a t t t t	5220
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g t a g a t t a t a	t a t t t t a g a t	t t t t t a a a t t	a g a g a t a t a t	t t a a t a t c g a	a g g t t a t t a a	5340
t g a g t t t t t a	t t t t t t g t t t	t t t a a t t g t a	a t g t t t t g a a	a g g t t t t t a g	t t t g g g t t t t	5400
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a t t t a a c g t t	t t t t g c g g t t	a g t t t t t t t a	a g a g g a t t t g	t t a t g t t t t t	a a g t t t g g t t	6600
a t t a g g t g t t	t c g g t t g g t t	t t a g g a t a g t	t t t g g t t a t t	c g t a a t t t t t	t a c g t a t t t t	6660
t t g t t t t t t t	g t t a g a g a t t	t t t t a t t t a g	t t a g a a g t g a	t a g a a a g g a a	t c g g a t g g t t	6720
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t t a t a t t t a t	g t a t t t t t t t	t t a a t t t a t t	t a g g t a t t t a	a t t a t t t a t t	t a t t t a t t t a	7380
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tt						8222

&lt;210&gt; 44

&lt;211&gt; 3051

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 44

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gcgttcgttt	tttttaatgt	atgttttagg	gggcgggttt	gcgcgggag	atggatacga	1860
ttggttttaa	agtttttttc	gtaaggtcgt	gggttggtata	gcgtggtgac	gtcgtaaacgc	1920
ggcgtagggt	gagagcgcg	gtttgcggac	gcgcgggtat	taaacgggtg	taggcgtagt	1980
agagtggtcg	ttgtttttt	aggttttagt	cggtcgtcgc	gacgttcgtt	cgttcgtttt	2040
gaggtttttg	aagtcgaaat	tagttagatt	tttttttttt	tcgtttgttt	gtagcggcgt	2100
tgttgttatt	tcgtttatt	gttcgagcgc	cgtttgtttt	agggttttat	ttttaagaag	2160
gtgttgagg	tatttaagga	ttttattaac	gaggtttgtt	gggatattag	ttttagcgg	2220
gtaaatttgt	agagtatgga	ttcgttttac	gttttttttg	tgtagtttat	tttgcggttt	2280
gaggggttcg	atatttatcg	ttgcgatcgt	aatttggtta	tgggcgtgaa	ttttattagg	2340
tgaagtttcg	ggtttcggga	agtcggtttc	ggttcgtttg	tattttcgtt	gtttgcccgg	2400

41

agcgttttcg	agtttagttt	ttattgggtg	gcgtgggtta	ttcgcgtttt	tttattgggt	2460
tgttacgtag	tggggtgggg	tttagttgag	cgcgcggttc	ggaaaagttc	gcgttggttg	2520
ttgcgcgaat	ttgttttttc	gcgttaaagt	tataaagcgg	gtggtggcgg	gaaaattaag	2580
ggtttttttcg	tagtggttagg	aatattgttt	taggggtttt	tgtttattaa	attttggttg	2640
ttttgaatgg	acgttttagt	tgtgggtttt	ttgtttttga	gacgggttcg	gtgtgtgtgt	2700
cgggttggtt	tttaattttt	gggtttaagc	gattttttcg	gtttagtcgc	ggttgatttt	2760
aaatgtttta	taatgttttt	gcgagaaatg	tggtagtgtt	ttattttatt	tagtggttagg	2820
agattgtttt	tatttagaag	ggatattgtt	ggtggtattt	tagtataaat	attggttagat	2880
gcgttttaaa	acgtttgtat	taataatggg	attttttagt	agttcgttta	ttttttatta	2940
gttttgagac	ggtttgatgg	gtgagagtgg	taattttttt	taatcgcggt	cgaaatatag	3000
tttttttagta	gacggcggtg	atttttaaagt	atgtgttttt	tgtttttttag	t	3051

&lt;210&gt; 45

&lt;211&gt; 3051

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 45

attagaagat	aggagatata	tgttttaaaa	ttaacgtcgt	ttggtgaagg	gttgtatttc	60
gaacgcggtt	agaaggggtt	attattttta	tttattaggt	cgtttttagaa	ttggtggagg	120
gtaagcggat	tggttgagga	tgttattatt	aatgtagacg	ttttgggacg	tatttggttag	180
tatttatatt	aaaataattat	tagtagtggt	ttttttggat	agaaataatt	ttttgttatt	240
aagtaggatg	ataggttggt	atatttttcg	taagggtatt	ataaagtatt	taaagttaat	300
cgcgattgag	tcgggaggat	cgtttgagtt	tagaagttgg	agattagttc	gggtaataata	360
tcgagatcgt	tttagaaata	agaaagtatt	agttaaagcg	tttattttaag	gttaatatagga	420
tttagtgagt	aaagagtttt	ggaatagtggt	ttttggtatt	gcggaaaaat	ttttgatttt	480
ttcgttatta	ttcgttttgt	gattttggcg	cgaaaaagta	ggttcgcgta	gtagtttagcg	540
cgggtttttt	cgaatcgcgc	gttttagttg	gttttatttt	attgcgtggt	agggttaatga	600
gaaggcgcgg	atggttttacg	ttagttaatg	agggttaggt	tcgaaagcgt	tttcgttaag	660
tatcgagggt	gtaggcgggt	cggggtcggg	ttttcggggg	cgcgagggtt	atttggtgag	720
gtttacgttt	atggttaggt	tgcggtcgta	gcggtagggt	tcgaagtttt	tagatcgtag	780
ggtgagttgt	attaaagaga	cgtagggacga	gtttatggtt	tgtaggttta	tatcgttgga	840
gttaatatatt	tagtaggttt	cgttgatgag	gtttttgagt	gtttttaata	tttttttgag	900
gatggagttt	tggattaggg	gcgtttcgaa	tatggtggcg	gagtggtaat	aacgtcgtta	960
taggtaggcg	ggaaggagga	aagtttagtt	ggtttcggtt	ttaggagttt	tagagcgagc	1020
gggcgaacgt	cgcgacgatc	ggttgagatt	tagaaaagata	acgattattt	tgttacgttt	1080
gtaatcgttt	aatgtcgtcg	cgttcgttaag	cgcgcggttt	tattttgcgt	cgcggtgcga	1140
cgttattacg	ttgttttagt	tacggttttg	cggggaagat	tttaggggta	atcgtgttta	1200
tgtttttcgc	gaggttcgtt	tttttagagta	tatatgggag	gaagcggggg	tagggcgggg	1260
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tttagttttt	tattacgtgg	gtagaaagtt	tttagttacg	aaagtgaag	tgaaatgtaa	1440
attaagttta	tcgggaggaa	aagttttttt	gtagtttgaa	gagagtatag	ttgttgtaaa	1500
tatgtatttt	attttttaat	tatagaattg	atgcgttcgt	agggtgtata	agataaagag	1560
gtgaattggt	ttgttaaat	tagtttttta	ttattttttt	aatatttttt	cgtagttttt	1620
tataattatt	taatcgttta	ttgtataaag	tttttaaagt	ttttttaaaa	gttcgaatat	1680
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tatatttaat	tttatgttac	gtatattttt	tattatatta	gtattttttt	aaaatatagt	1800
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tatttaggta	attttggtga	tattaatttt	ttgaagtaat	atagttgagt	taaagagcgt	1920
atatatttaa	tatttttttt	tttttttttt	tttttttttg	agatggagtt	ttgttttggt	1980
atttaggttg	gagtatagtg	gcgcgatttt	agtttattgt	aattttcgtt	tttttaggtt	2040
aagtgagttt	ttttgttttag	tttttcogagt	atttgggatt	ataggtgttt	attattattt	2100
tcggttaatt	tttgattatt	tattattatt	tttttttagt	agagacgggg	ttttattatg	2160
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gtgttgggat	tataggcgtg	agttattacg	tttagttaga	aatttttaat	tgttatagga	2280
tggataggat	ttgttgtttt	ttttttgttt	tttttatatt	attttttttt	agttattttt	2340
aatatttttg	ttagtagaat	tttttagttt	ttttattatg	tatttaggta	tggagttagt	2400
tttatttggt	ttgttgaggg	aattgaagat	ttattttaag	tttgatatat	ttttaattat	2460



42

tatgggtttta	aatatTTTTT	TTTTTtagtt	tattttttat	ttttggaatg	TTTTTTTTTT	2520
TTTTTTTTTT	TTTTTTTTTT	TTTTTtagtt	tagttgtgat	tttatataag	gttttaagat	2580
gagttttata	gttttttttt	tggattttgt	ggagatttaa	ttttatTTTT	ttgtattagt	2640
tttttaaagt	attggggttag	ttttaaaagg	gaataggaaa	gggtatTTTT	attataagat	2700
gtttttaatt	gtaatatgta	ttattatTTT	atgtagtata	agtaagaaaa	ttgttaaatt	2760
TTTTTTTTTT	ttatTTTTTT	gagaggggag	TTTTTTTTTT	atatttaggt	tggagtgtag	2820
tggtaaagatg	attatTTTatt	gttattTTTT	atttttaggt	tgcagagatt	TTTTtgTTat	2880
TTTTtaaata	gttgggatta	taggggtgta	ttattgtatt	tggttaattt	ttaaattgtt	2940
ttgtagagag	aaggTTTTat	tatgtcgttt	tggttggttt	ttaattTTTT	ggtttaagta	3000
atttttttaa	tcggtTTTTT	aaagtgtgag	attataggta	tgagttattg	t	3051

&lt;210&gt; 46

&lt;211&gt; 3664

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 46

gaatttttaa	ttattatTTT	gtttttttat	ttttaatatt	tttagcggta	attgtatTTT	60
atttatTTTT	gtgtttatga	agtatatgt	tgtatTTTgg	agatattata	tagaagttgg	120
tttaaagggt	aataaattgt	atttgTTaag	atgatagtat	ggatttataa	taatttttga	180
aatTTtataa	agaaagatga	ttgagTTaaa	ttgatataatt	tagtattata	gttTgtggaag	240
ttTgtggTTT	TTTTtatgtt	aaaggTaaaa	tggaataaat	gaattagTTg	ttatTTtttag	300
atgatataga	agttgaattt	TTTTtaagtt	TTTTaaagag	aaagatatat	taaaattttat	360
taaaattagt	attataaaatt	gtatagaata	gtttggTTTT	ttagagtatt	TTTTtggggT	420
agtTggtTTT	tattttaaaaa	atgaagTTTg	gattTTtaaat	ataagTTTta	aaagtTatta	480
ggTTtttagta	ggagTTttta	aattTTttatt	tatattatga	TTTTtaaatg	TTTTgtTTTT	540
gttGtttagg	aataaaaaatt	gttaaggTTg	agtttagatgt	aatgtataat	atttagatgat	600
agatgtagtT	tattgtgtta	aaaggTgtta	aatattTTaag	tgtatgtgat	atattTTtata	660
ttataagggT	aaggaagTTT	tattTTtata	tttgattTTg	gtttTTtaaat	atatattTTgt	720
agtggatgtg	gtaattagtT	tttGttttat	gttttatttag	gattaatgta	atttagaaatt	780
tttaaaatgg	atttttatta	tgttttagttt	tgttttattg	ttttattTgt	ttattTTtaaa	840
aagattTTtag	aatggTgttt	aggTtgggtt	ggTggTttac	gtttgttaatt	ttagtatttt	900
gggaggttaa	ggtaggagga	ttgttggagt	ttaggagTTg	gagattagtT	agggtaatat	960
aaggagatat	tgttttcgta	aataaataga	aaaaaaaaaa	gtaagaaaga	aagagagaaa	1020
gaaggaaaaga	ataagaaaaa	ggaaaaaaaa	attaattgta	gtaagtgtag	aaattTTTTT	1080
taaggatatg	atggatggta	atgaagatag	agtttttgat	attggattat	ttaaagtata	1140
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agttaggcgt	ggtggcgagt	gtttgtagtc	tgaattTcgt	tggaggtTga	ggTcgattgt	1260
ttgagtttag	gatttgagg	ttagtatcgc	taataataatg	agatttagtt	tttaaatgta	1320
tgtttttttt	tatatattta	aaattttgat	gtgaaaatat	tttaaaattt	aatatatTTT	1380
aaatgtTTTt	aattgtataa	taaataaaat	gtaaataata	aaataattta	atattaaatt	1440
taaaaatgag	gtagaaataa	agtatagcga	tataaataat	aaattTTTTT	ttatatTTTT	1500
gaggcggttt	tttgagtttt	ttatTTTTTT	tttaaggTTa	ttgaaatgtg	TTTTttggag	1560
ttagtTcgtA	aattacgtat	ttagaaaaat	ataattatat	attTTtaatt	ttagtattta	1620
gaagtgaag	taatggaatt	tcgatgtaaa	tataatatta	TTTTtttgat	gagttatttt	1680
gagtataata	aatttgaatt	gtgttaatgt	tgggagaaaa	aattttaaag	aagaacggag	1740
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ggagagcgTt	tgcgttcgta	tttagttggc	gttcgtTTTT	ttgtttTTTT	tttagtcgTt	1860
TTTTTTTTTT	TTTTtcgcgt	tttagttatt	cgggaaggTt	tgttttagcgt	agttgggttt	1920
tgattggTtg	TTTTgaaagt	ttacgggtta	ttcgattggT	gaattcgggg	TTTTtttagcg	1980
cggTgagTtt	gaaattgttc	gtatttggtt	ttaaagTtgg	TTTTttggaaa	ttgagcggag	2040
agcgacgcgg	ttgttgtagt	tgtcgtttcg	gtcgtcgcgg	aataataagt	cgggtatagt	2100
ggttggggTt	aggTcgtgt	ttaggggacg	gtcgaaggTt	tcggagggcg	agtattgagg	2160
aacggggTtt	tttaagaagg	tcggattTga	ggtttagggat	ttgcgcgggg	ttcggTtggg	2220
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ggtttttagag	ggaggttatg	ggagtttagt	ttggcggggT	taggcggcgt	gggggtgggg	2340
gttcggggTt	gtagttatga	gtcgaaggTc	gttgtttTta	ggaattgttt	gaagaggttt	2400
tcgggtttTg	attgaagtaa	aattcgtttt	tttacgtTga	gcgtttTcgg	agtagtaggt	2460
TTTTTTTTTg	gttcggTttt	tgcgcgtttt	gttatTtcgt	ttcgggggtg	ggaaagtggg	2520

43

atcgtattttg	ggttaggtat	aggcgcgggt	tttttagagtt	aggtcgtttt	gagaaatcga	2580
tttttgaatt	cgtgggtttt	tttgggtttg	ttttgtttac	ggtttcggtt	tatttgaggt	2640
tgtaagaat	agttagttta	ggtattttaat	ttttatgagt	agtgtagtgt	agtttttttt	2700
atttaaaaag	atataattttt	gttttttttt	ggattgattt	ttttttgaag	atgaatgtga	2760
gaaatagaat	ttaaagggtt	attttgagtg	tgttttataa	atgattttat	tattagtgtat	2820
gggtgttaga	aaaatttagg	taattatgtt	tggagttttg	acgtttttatt	ggtaaattttt	2880
agaggaagtt	ttgaattttt	ggttggttagt	tttttttttt	aattttattta	ttgtgattta	2940
agtatatcgt	ttttttgttt	agtttttgat	attgtttagg	ttataattag	aattttatttt	3000
gtggaattgt	atttttttatt	ttttttgttt	acggtgttta	agtataattt	tttattttttg	3060
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agattaaaat	tgttacgtgg	gtaattgtaa	atatttaaaa	tgtttttgtc	gttttaaaaa	3180
aaattttttt	aatgaatagt	ttgaatatat	tttagttttg	ttttttataat	tttaaaaaaat	3240
aatgttttga	ataaaattag	gagtattata	aagcgggtata	aatttggtga	aataatttggg	3300
tataattttg	gtatacgtag	gtattatttt	aaagtgttat	aaaaagggaa	tataatttttg	3360
gtttttgttt	ggaaagaata	gataagttga	tatttgaatt	tgaggggttat	tttgatttga	3420
agaagttaaa	tattatatta	gtatagaata	tatacgtttt	taatgtttta	ggagttaaga	3480
aggatggaaa	gttattgtag	gagtttttat	ttttgtaaat	attgtttgaa	tgtaattaat	3540
atztatggaa	tatttatgtt	gtagtaggta	ttgtggaaaa	tgttttatat	atagtgtttt	3600
attaatgttt	aaaattattc	gagtttagtg	tatgtaatta	agttagtttt	ttttttattt	3660
tagg						3664

&lt;210&gt; 47

&lt;211&gt; 3664

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 47

tttaaagtga	aggaaaaaatt	gatttaatta	tatgttatta	attcgagtag	ttttaagtat	60
taatgaaata	ttatatgtaa	agtatttttt	atagtgttta	ttgtagtata	agtattttat	120
aaatattaat	tatatttaaa	taatatttat	aagggttaagg	atttttataa	taatttttta	180
ttttttttga	tttttgagat	attggaaacg	tgtatatatt	atattagtgt	agtatttagt	240
tttttttaat	taaaataaatt	tttaaattta	ggtattaatt	tgttttattt	ttttaataaa	300
aaattagaat	tatgtttttt	ttttgtgata	ttttaaagta	gtatttgcgt	gtgttaaagt	360
tatatttaaa	tatttttaata	aatttatatc	gttttatgat	atttttaatt	ttatttaaaa	420
tattattttt	taaagttatg	aggatagaat	taaaatgtgt	ttagattgtt	tattaggaaa	480
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ttatagataa	aatttttaatt	ataattttaa	taatgttaag	gattaagtag	aaaaacggta	720
tatttgaatt	ataatgaatg	ggttaaaaaa	aaaagttgat	aattagaagt	ttaagatttt	780
ttttgaagtt	tgttaatgaa	acgttagaat	tttaggtata	attattttaga	tttttttaaat	840
atttattatt	gatgataaaa	ttattttata	agtatattta	gaatagattt	ttaagtttta	900
ttttttatat	ttatttttta	agaaaaatta	atthagaaag	aagtagagat	gtattttttt	960
aaatgaaaaa	aattgtatta	tattgtttat	gagagttaag	tgtttggttt	gattgttttt	1020
aatagtttta	ggtgggtcgg	ggtcgtgggt	agagtaagtt	taagaaggtt	tacgagtttt	1080
aggatcgggt	ttttagggcg	gtttgggttt	agggttcgc	gtttgtattt	ggtttaggtg	1140
cggttttatt	tttttatttt	cgaggcggga	tggtagggcg	cgtaggggtc	gggttttagga	1200
aagagtttgt	tatttcgtag	gcgttttagc	tggtgggagc	gattttgttt	taattaagat	1260
tcgaaagttt	ttttaggtag	tttttgagaa	tagtcgattt	cgatttatga	ttgtagtttc	1320
gaatttttat	ttttacgtcg	tttaatttcg	ttaggttggg	tttttatagt	ttttttttaa	1380
agttagttta	gtgtggttta	tacggttaga	tttaggcgaa	ggttttgttt	atcgtttttc	1440
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aaagggcggg	tagagaaaaa	gtaggagggc	gggcgttaat	tgagtgcgag	cgtaagcgtt	1860
tttttttagt	cgggagagtg	tcgtttttatt	gttttttagt	agcggagtag	gaagttattg	1920

ttcgtttcgt	ttttttttta	aatttttttt	tttagtattg	gtatagttta	aattttattat	1980
atttaaaata	gtttattaaa	aaagtgatat	tgtgtttata	tcgagatttt	attatttttta	2040
tttttaatat	ttagggttag	gagtgtatag	ttatgttttt	ttaaatgcgt	gatttgcggg	2100
ttggttttta	ggagtatatt	ttagtatttt	taagaaggaa	atggaaaatt	taaaagatcg	2160
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atttgaaatg	tattaaattt	taaaatattt	ttatattaga	attttaaata	tatagagaga	2340
ggtatgtatt	tagagattgg	gttttattat	gttgcgtatg	ttggttttta	aattttgggt	2400
ttaagtaatc	ggtttttagt	tttagcgtag	ttgcgattat	aggatttcgt	tattacgttt	2460
ggtttaata	taattaaaat	ataaaaaatt	ataattttat	gaggggagag	aaagagggtt	2520
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tttgagagag	atttttgtat	ttattgtagt	taattttttt	tttttttttt	ttgttttttt	2640
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taaggatttt	tggttgtatt	gattttaatg	ggatatagaa	taaagattga	ttattatatt	2940
tattataagt	atgtattaga	agatttaaat	taggatattg	aatggaattt	ttttattttt	3000
gtgatataga	gtatattata	tgtatttaag	tgtttaatat	tttttaatat	aatgagttgt	3060
atttattatt	tggtgttata	tattgtattt	ggtttaattt	tgataatttt	tggttttagg	3120
taatagaggt	agagtatttg	ggagtattgg	tgtaggtagg	aatttaagaa	ttttgttggt	3180
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tattgtttta	gagaagtgtt	ttagggaatt	agggtgtttt	gtgtagttta	tggtattggg	3300
tttaatgaat	tttaatatgt	tttttttttt	gagaggttta	agaggaattt	aatttttata	3360
ttattttaaa	atggttaatta	atttattata	tttattttat	ttttaatata	aagaagatta	3420
taagttttta	tagttgtgat	gttaggtgta	ttaatttaat	ttaattattt	ttttttataa	3480
agtttttagag	gttattgtaa	atttatgtta	ttatttggg	aaatatagtt	tattattttt	3540
tgggttaatt	tttatgtagt	atttttaaa	tgtaatatgt	tgttttatag	atataaaaa	3600
gaataagata	tagttatcgt	tggaggtgtt	gagggtaggg	gagtaggatg	ataattggga	3660
at						3664

&lt;210&gt; 48

&lt;211&gt; 1729

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 48

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aggtttagtt	aaatttgaat	tttagataaa	taatgaataa	tttgttagta	taaatatggt	120
ttatgtaata	ttttgttgaa	attaaaaaaa	aaaaaaaaag	tttttttttt	attttttatt	180
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45

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acggtagcgg	ttcagatgg	tttatttaag	agattggcgt	tttttaggt		1729

&lt;210&gt; 49

&lt;211&gt; 1729

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 49

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&lt;210&gt; 50

&lt;211&gt; 12963

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 50

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49

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&lt;210&gt; 51

&lt;211&gt; 12963

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 51

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55

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tattattttaa	ttatttttta	taaggtttta	tttttaatat	tggggtttat	aatttgatat	3060
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tttttgttat	ttgttttgag					3500

&lt;210&gt; 54

&lt;211&gt; 3315

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 54

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56

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gttttagtaga	gttttattta	tttttttaat	gtttttgttt	aatgtgtttt	ttaaattttt	3300
ttttatttag	attat					3315

&lt;210&gt; 55

&lt;211&gt; 3315

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 55

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gttttgggtt	ttttgtgtgt	atttttgggt	tgtgtgtttt	gtggtttttg	tagtttttta	420
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57

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ttatttaatt	tttataaaaa	tagattattg	ttggataata	tgtaaatgta	gttgaagtta	2640
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gtggggagtg	agggg					3315

&lt;210&gt; 56

&lt;211&gt; 7369

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 56

tagaagtttt	tttttagagt	gtgtttgggt	atataatttaa	gtgtatgggt	gtaaattttt	60
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aattattttta	ttgtgaagtt	tagtattata	aaaatttttaa	taattttatta	taagtttttta	240
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59

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&lt;210&gt; 57

&lt;211&gt; 7369

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 57

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&lt;210&gt; 58

&lt;211&gt; 2986

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 58

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tgttgtgttt	atTTTTggtt	aatTTTTaaa	agaaagatgt	ttgttttggg	ttttttttta	900
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taataaagat	tgtgtgtttt	ttttaaaaat	tttttttgtg	tttaga		2986

&lt;210&gt; 59

&lt;211&gt; 2986

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 59

tttaagtgt	ggggaaattt	ttaaaaagaa	tatataattt	ttattagaaa	aaataatggt	60
tggtagaagt	tgggttttata	gttaggggat	gaaaaattta	ttatttttat	ttataatttt	120
ggatttggt	tttttagtgg	tttttaagg	tataaaatga	ggttgttttt	aaaaaataga	180
aagtgtattt	tttgttttag	agtttttttt	ttagtttagga	ggtggattgt	tgataaatat	240
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aagtgaatgt	ttttttttta	gaaattagtt	aggagtagat	gtggtattta	ttatgaattt	2160
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atagatgagt	ttgatgtgta	gtttgttttt	ttagtataat	tttgtttatt	ttgtgtgttt	2940
atttttattt	atatgtttta	gaatgtgata	gttggaaagg	attttt		2986

&lt;210&gt; 60

&lt;211&gt; 5666

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 60

aaaattagaa	tttttatttt	tttgtgtttg	ttatattttt	tagtgtttgt	taattttttt	60
ttgtaagtga	gggtggtgga	gggtgtttat	aattttttta	gggagtaagt	tttttttggg	120
tttttttttt	tttttttttt	tttttttttt	tgagattaag	ttttgttttt	gttttttagg	180
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65

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ttgattattt	tgtttttttt	ggtagg				5666

&lt;210&gt; 61

&lt;211&gt; 5666

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 61

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tgtaaggagg	taaaagtttt	aattttt				5666

&lt;210&gt; 62

&lt;211&gt; 5085

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 62

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68

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&lt;210&gt; 63

&lt;211&gt; 5085

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 63

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&lt;210&gt; 64

&lt;211&gt; 8222

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 64

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72

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&lt;210&gt; 65

&lt;211&gt; 8222

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 65

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&lt;210&gt; 66

&lt;211&gt; 3051

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 66

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&lt;210&gt; 67

&lt;211&gt; 3051

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 67

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76

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&lt;210&gt; 68

&lt;211&gt; 3664

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 68

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tagg						3664

&lt;210&gt; 69

&lt;211&gt; 3664

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 69

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<211> 1729  
 <212> DNA  
 <213> Artificial Sequence

<220>  
 <223> chemically treated genomic DNA (Homo sapiens)

<400> 70

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<210> 71  
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 <212> DNA  
 <213> Artificial Sequence

<220>  
 <223> chemically treated genomic DNA (Homo sapiens)

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80

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&lt;210&gt; 72

&lt;211&gt; 12963

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 72

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&lt;210&gt; 73

&lt;211&gt; 12963

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 73

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&lt;210&gt; 74

&lt;211&gt; 3500

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 74

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88

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&lt;210&gt; 75

&lt;211&gt; 3500

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; chemically treated genomic DNA (Homo sapiens)

&lt;400&gt; 75

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89

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&lt;210&gt; 76

&lt;211&gt; 20

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Detection primer for ESR1

&lt;400&gt; 76

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&lt;210&gt; 77

&lt;211&gt; 22

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence



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&lt;223&gt; Detection oligonucleotide for MB

&lt;400&gt; 325

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&lt;400&gt; 326

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&lt;223&gt; Detection oligonucleotide for CEA

&lt;400&gt; 522

aatagatacg gagaggga

18

&lt;210&gt; 523

&lt;211&gt; 18

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Detection oligonucleotide for CEA

&lt;400&gt; 523

aatagatatg gagaggga

18

&lt;210&gt; 524

&lt;211&gt; 20

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Detection oligonucleotide for CEA

&lt;400&gt; 524

tggttaaagt tgtgggagat

20

&lt;210&gt; 525

&lt;211&gt; 20

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Detection primer for CEA

&lt;400&gt; 525

tcctgagtga tgtctgtgtg

20

&lt;210&gt; 526

&lt;211&gt; 20

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Detection primer för p16

176

&lt;400&gt; 526

atgacaccaa acaccccgat

20

&lt;210&gt; 527

&lt;211&gt; 19

&lt;212&gt; DNA

&lt;213&gt; Artificial Sequence

&lt;220&gt;

&lt;223&gt; Detection primer for p16

&lt;400&gt; 527

ctgtccctca aatcctctg

19





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(81) Designated States (*national*): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.

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**WO 03/014388 A3**

(54) Title: **METHOD AND NUCLEIC ACIDS FOR THE ANALYSIS OF COLON CANCER**

(57) Abstract: The present invention relates to chemically modified genomic sequences, oligonucleotides and/or PNA-oligomers for detecting the cytosine methylation state of genomic DNA, as well as to methods for ascertaining genetic and/or epigenetic parameters of genes for use in the characterisation, grading, staging, and/or diagnosis of colon cancer, or the predisposition to colon cancer.

# INTERNATIONAL SEARCH REPORT

International Application No  
PCT/EP 02/08939

A. CLASSIFICATION OF SUBJECT MATTER  
IPC 7 C12Q1/68

According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)  
IPC 7 C12Q

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data, BIOSIS, MEDLINE, SEQUENCE SEARCH

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	TOYOTA MINORU ET AL: "Identification of differentially methylated sequences in colorectal cancer by methylated CpG island amplification" CANCER RESEARCH, AMERICAN ASSOCIATION FOR CANCER RESEARCH, BALTIMORE, MD, US, vol. 59, no. 10, 15 May 1999 (1999-05-15), pages 2307-2312, XP002211911 ISSN: 0008-5472 page 2307 -page 2312; figure 1 ---	1
X	TOYOTA MINORU ET AL: "Methylation profiling in acute myeloid leukemia." BLOOD, vol. 97, no. 9, 1 May 2001 (2001-05-01), pages 2823-2829, XP002226261 ISSN: 0006-4971 page 2823 -page 2829; table 1 ---	1-40
-/--		

☒ Further documents are listed in the continuation of box C.

☒ Patent family members are listed in annex.

### \* Special categories of cited documents :

"A" document defining the general state of the art which is not considered to be of particular relevance

"E" earlier document but published on or after the international filing date

"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

"O" document referring to an oral disclosure, use, exhibition or other means

"P" document published prior to the international filing date but later than the priority date claimed

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"&" document member of the same patent family

Date of the actual completion of the international search

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Date of mailing of the international search report

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Name and mailing address of the ISA

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## INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 02/08939

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT		
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	LI LONG-CHENG ET AL: "Frequent methylation of estrogen receptor in prostate cancer: Correlation with tumor progression." CANCER RESEARCH, vol. 60, no. 3, 1 February 2000 (2000-02-01), pages 702-706, XP002226262 ISSN: 0008-5472 the whole document ---	1-40
A	WO 00 44934 A (OLEK ALEXANDER ;EPIGENOMICS GMBH (DE)) 3 August 2000 (2000-08-03) the whole document ---	1-40
A	WO 99 28498 A (OLEK ALEXANDER ;WALTER JOERN (DE); EPIGENOMICS GMBH (DE); OLEK SVE) 10 June 1999 (1999-06-10) the whole document ---	1-40
A	REIN ET AL: "Identifying 5-methylcytosine and related modifications in DNA genomes" NUCLEIC ACIDS RESEARCH, OXFORD UNIVERSITY PRESS, SURREY, GB, vol. 26, no. 10, 1998, pages 2255-2264, XP002143106 ISSN: 0305-1048 page 2307 -page 2312; figure 1 ---	17-21
A	TOYOTA MINORU ET AL: "CpG island methylator phenotypes in aging and cancer." SEMINARS IN CANCER BIOLOGY, vol. 9, no. 5, October 1999 (1999-10), pages 349-357, XP002226263 ISSN: 1044-579X page 349 -page 257; figure 1 ---	1-40
A	BAYLIN S B ET AL: "DNA hypermethylation in tumorigenesis: epigenetics joins genetics" TRENDS IN GENETICS, ELSEVIER, AMSTERDAM, NL, vol. 16, no. 4, April 2000 (2000-04), pages 168-174, XP004194021 ISSN: 0168-9525 the whole document -----	1-40

# INTERNATIONAL SEARCH REPORT

International application No.  
PCT/EP 02/08939

## Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. ☐ Claims Nos.:  
because they relate to subject matter not required to be searched by this Authority, namely:
2. ☐ Claims Nos.:  
because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3. ☐ Claims Nos.:  
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

## Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

see additional sheet

1. ☐ As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. ☐ As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. ☐ As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. ☒ No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

1-40 IN PART

Remark on Protest

- ☐ The additional search fees were accompanied by the applicant's protest.
- ☐ No protest accompanied the payment of additional search fees.

## FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-40 (in part)

A method to determine the methylation status of CpG dinucleotides in the estrogen receptor gene for the diagnosis of colon cell proliferative disorders.

(i) Said method, involving bisulfite-PCR of pretreated genomic DNAs (SEQ ID NO: 33,33,54,55) with specific primers (SEQ ID NO: 76, 77) using labelled nucleotides, and involving detection by hybridisation to an array of probes (SEQ ID NO: 98-145) and relevant isolated nucleic acids.

(ii) Said method to determine the methylation status of CpG dinucleotides in the estrogen receptor gene, making use of methylation-sensitive restriction enzymes and corresponding detection primers (SEQ ID NO: 12-18).

2. Claims: 1-40 (in part)

Idem as subject matter 1, but limited to the p21 gene, wherein:  
pretreated genomic sequences are SEQ ID NO: 34,35,56,57  
primers for amplification are SEQ ID NO: 78,79  
probes for detection are SEQ ID NO: 146-169

3. Claims: 1-40 (in part)

Idem as subject matter 1, but limited to the p27 gene, wherein:  
pretreated genomic sequences are SEQ ID NO: 36,37,56,57  
primers for amplification are SEQ ID NO: 80, 81  
probes for detection are SEQ ID NO: 170-201

4. Claims: 1-40 (in part)

Idem as subject matter 1, but limited to the p16 gene, wherein:  
pretreated genomic sequences are SEQ ID NO: 38,39,60,61  
primers for amplification are SEQ ID NO: 82, 83  
probes for detection are SEQ ID NO: 202-249

5. Claims: 1-40 (in part)

Idem as subject matter 1, but limited to the progesterone receptor gene, wherein:  
pretreated genomic sequences are SEQ ID NO: 40,41,62,63  
primers for amplification are SEQ ID NO: 84, 85  
probes for detection are SEQ ID NO: 250-313

## FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

## 6. Claims: 1-40 (in part)

Idem as subject matter 1, but limited to the myoglobin gene,  
wherein:  
pretreated genomic sequences are SEQ ID NO: 42,43,64,65  
primers for amplification are SEQ ID NO: 86, 87  
probes for detection are SEQ ID NO: 313-357

## 7. Claims: 1-40 (in part)

Idem as subject matter 1, but limited to the pcna gene,  
wherein:  
pretreated genomic sequences are SEQ ID NO: 44,45,66,67  
primers for amplification are SEQ ID NO: 88, 89  
probes for detection are SEQ ID NO: 358-405

## 8. Claims: 1-40 (in part)

Idem as subject matter 1, but limited to the cdc2 gene,  
wherein:  
pretreated genomic sequences are SEQ ID NO: 46,47,68,69  
primers for amplification are SEQ ID NO: 90, 91  
probes for detection are SEQ ID NO: 406-435

## 9. Claims: 1-40 (in part)

Idem as subject matter 1, but limited to the c-erbB2 gene,  
wherein:  
pretreated genomic sequences are SEQ ID NO: 48,49,70,71  
primers for amplification are SEQ ID NO: 92, 93  
probes for detection are SEQ ID NO: 436-451 and  
detection primers are SEQ ID NO: 22 and 23.

## 10. Claims: 1-40 (in part)

Idem as subject matter 1, but limited to the p53 gene,  
wherein:  
pretreated genomic sequences are SEQ ID NO: 50,51,72,73  
primers for amplification are SEQ ID NO: 94, 95  
probes for detection are SEQ ID NO: 452-489 and  
detection primers are SEQ ID NO: 24-27.

## 11. Claims: 1-40 (in part)

Idem as subject matter 1, but limited to the CEA gene,  
wherein:  
pretreated genomic sequences are SEQ ID NO: 52,53,74,75  
primers for amplification are SEQ ID NO: 96, 97

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

probes for detection are SEQ ID NO: 490-523 and  
detection primers are SEQ ID NO: 28-31.

**INTERNATIONAL SEARCH REPORT**  
Information on patent family members

International Application No  
**PCT/EP 02/08939**

Patent document cited in search report		Publication date	Patent family member(s)	Publication date
WO 0044934	A	03-08-2000	DE 19905082 C1	18-05-2000
			AU 3144700 A	18-08-2000
			CA 2359182 A1	03-08-2000
			WO 0044934 A2	03-08-2000
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WO 9928498	A	10-06-1999	DE 19754482 A1	01-07-1999
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			PT 1034309 T	31-10-2002
			SI 1034309 T1	31-08-2002
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